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INTERNATIONAL APPLICATION PUBLISH	HED U	IDER THE PATENT COOPERATION	N TREATY (PC1)
51) International Patent Classification 6:		11) International Publication Number:	WO 99/13886
A61K 31/70, 48/00, C07H 21/00, 21/04, C12N 5/10	A1	43) International Publication Date:	25 March 1999 (25.03.99)
21) International Application Number: PCT/US (22) International Filing Date: 17 September 1998 ((30) Priority Data: 60/059,160 09/093,972 9 June 1998 (09.06.98) (71) Applicant (for all designated States except US): EA OLINA UNIVERSITY [US/US]; 210 Spilman Greenville, NC 27858-4353 (US). (72) Inventor; and (75) Inventor/Applicant (for US only): NYCE, Jon [US/US]; 59 Sayre Drive, Princeton, NJ 08540 ((74) Agents: AMZEL, Viviana et al.; Pretty, Scl Poplawski, 444 South Flower Street, Los An 90071 (US).	(17.09.9 .97) U AST CA Building mathan, (US).	GH, GM, HR, HU, ID, IL, IS, LC, LK, LR, LS, LT, LU, LV, MX, NO, NZ, PL, PT, RO, RI TJ, TM, TR, TT, UA, UG, US, patent (GH, GM, KE, LS, MW, patent (AM, AZ, BY, KG, KZ, I patent (AT, BE, CH, CY, DE, IE, IT, LU, MC, NL, PT, SE) CG, CI, CM, GA, GN, GW, M Published With international search report Before the expiration of the claims and to be republished amendments.	JP, KE, KG, KP, KR, KZ, MD, MG, MK, MN, MW, J, SD, SE, SG, SI, SK, SL, UZ, VN, YU, ZW, ARIPO SD, SZ, UG, ZW), Eurasian MD, RU, TJ, TM), European DK, ES, FI, FR, GB, GR, OAPI patent (BF, BJ, CF, L, MR, NE, SN, TD, TG).
(54) Title: MULTIPLE TARGET HYBRIDIZING NUKITS AND APPLICATIONS (57) Abstract Antisense oligonucleotides which bind to two or the second sec			SITIONS, FORMULATION

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MULTIPLE TARGET HYBRIDIZING NUCLEIC ACIDS, THEIR PREPARATION, COMPOSITIONS, FORMULATION, KITS & APPLICATIONS

BACKGROUND OF THE INVENTION

Field of the Invention

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This invention relates to multiple target anti-sense oligonucleotides (MTA oligos) of low or no adenosine content. The present agents are effective in the prophylaxis and treatment of diseases and conditions associated with inflammation, impaired airways, including lung disease and diseases whose secondary effects afflict the lungs of a subject. The agents are targeted to specific genes, genomic flanking regions, initiation codon, intron-exon borders, and the like, or the coding and non-coding regions of RNAs, including those encoding certain proteins, particularly those associated with diseases having multiple mediators, by affecting (either attenuating or enhancing) various contributing pathways. Examples are pulmonary diseases such as allergies, asthma, impeded respiration, pain, cystic fibrosis and cancers such as leukemias, e.g. colon cancer, and the like. The present agent may easily be administered prophylactically and therapeutically in conjunction with other therapies, or may be utilized as a substitute for therapies that have significant, negative side effects.

Description of the Background

Respiratory ailments, associated with a variety of diseases and conditions, are extremely common in the general population, and more so in certain ethnic groups, such as African Americans. In some cases they are accompanied by inflammation, which aggravates the condition of the lungs. Asthma, for example, is one of the most common diseases in industrialized countries. In the United States it accounts for about 1% of all health care costs. An alarming increase in both the prevalence and mortality of asthma over the past decade has been reported, and asthma is predicted to be the preeminent occupational lung disease in the next decade. While the increasing mortality of asthma in industrialized countries could be attributable to the increased reliance upon beta agonists in the treatment of this disease, the underlying causes of asthma remain poorly understood.

Anti-sense oligonucleotides have received considerable theoretical consideration as potential useful pharmacological agents in human disease. Their practical application in actual models of human disease, however, has been somewhat elusive. One important impediment to their effective application has been a difficulty in finding an appropriate route of administration to deliver them to their site of action. Many in vivo experiments were conducted by administering anti-sense oligonucleotides directly to specific regions of the brain. These applications, however, necessarily have limited clinical utility due to their invasive nature.

The systemic administration of anti-sense oligonucleotides also presents significant problems, not the least being an inherent difficulty in targeting disease-involved tissues. In

contrast, the lung is an excellent target for the direct administration of anti-sense oligonucleotides, and provides a non-invasive and a tissue-specific route. The delivery of anti-sense agents to the lung has been relatively undeveloped.

Adenosine may constitute an important mediator in the lung for various diseases, including bronchial asthma. Its potential role was suggested by the finding that asthmatics respond favorably to aerosolized adenosine with marked bronchoconstriction whereas normal individuals do not. An asthmatic rabbit animal model, the dust mite allergic rabbit model for human asthma, responded in a similar fashion to aerosolized adenosine with marked bronchoconstriction whereas non-asthmatic rabbits showed no response. More recent work with this animal model suggested that adenosine-induced bronchoconstriction and bronchial hyperresponsiveness in asthma may be mediated primarily through the stimulation of adenosine receptors. Adenosine has also been shown to cause adverse effects, including death, when administered therapeutically for other diseases and conditions in subjects with previously undiagnosed hyper reactive airways.

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A handful of medicaments have been available for the treatment of respiratory diseases and conditions, although in general they all have limitations. Theophylline, an important drug in the treatment of asthma, is a known adenosine receptor antagonist which was reported to eliminate adenosine-mediated bronchoconstriction in asthmatic rabbits. A selective adenosine A₁ receptor antagonist, 8-cyclopentyl-1, 3-dipropylxanthine (DPCPX) was also reported to inhibit adenosine-mediated bronchoconstriction and bronchial hyperresponsiveness in allergic rabbits. The therapeutic and preventative applications of currently available adenosine A₁ receptor-specific antagonists are, nevertheless, limited by their toxicity. Theophylline, for example, has been widely used in the treatment of asthma, but is associated with frequent, significant toxicity resulting from its narrow therapeutic dose range. DPCPX is far too toxic to be useful clinically. The fact that, despite decades of extensive research, no specific adenosine receptor antagonist is available for clinical use attests to the general toxicity of these agents.

Anti-sense oligonucleotides have received considerable theoretical consideration for their potential use as pharmacological agents in human disease. Finding practical and effective applications of these agents in actual models of human disease, however, have been few and far between, particularly because they had to be administered in large doses. Another important consideration in the pharmacologic application of these molecules is their route of administration. Many in vivo applications have involved the direct administration of anti-sense oligonucleotides to limited regions of the brain. Such applications, however, have limited clinical utility due to their invasive nature.

The systemic administration of anti-sense oligonucleotides as pharmacological agents

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has been found to have also significant problems, not the least of which being a difficulty in targeting disease-involved tissues. That is, the necessary dilution of the anti-sense oligonucleotide in the circulatory system makes extremely difficult to attain a therapeutic dose at the target tissue by intravenous or oral administration. The bioavailability of orally administered anti-sense oligonucleotides is very low, of the order of less than about 5%.

Anti-sense oligonucleotides have been used in therapy by many, including the present inventor, who in his previous work successfully treated various diseases and conditions by direct administration of these agents to the lung. In many instances, other workers have had to face the difficulties associated with the delivery of DNA molecules to a desired target. Thus, the route of administration may be of extreme importance for treating generalized diseases and conditions as well as those which are localized.

Accordingly, there is a need for effective ways to implement anti-sense therapy, particularly those which are highly effective for the treatment or diseases and conditions which are of a complex nature. Examples of the latter are those diseases and conditions which are associated with complex pathways and multiple endogenous agents which either alone, or in combination with one another, result in pathological conditions to the subject.

SUMMARY OF THE INVENTION

This invention relates to an agent, comprising an anti-sense oligonucleotide hybridizing to two or more mRNAs, or multiple target anti-sense oligos (MTA oligos), such as those that correspond to target genes, genomic flanking regions, the initiation codon, intron-exon borders, and the like, and the entire sequence of RNAs, including the coding region of mRNA and non-coding RNA segments such as the 5' cap or end and the 3' end, e.g. poly-A segment, and RNAs encoding proteins known to be associated with one or more diseases or conditions and mixtures thereof. The mRNAs, for example, may encode polypeptide(s) such as transcription factors, stimulating and activating factors, interleukins, interleukin receptors, chemokines, chemokine receptors, endogenously produced specific and non-specific enzymes, immunoglobulins, antibody receptors, central nervous system (CNS) and peripheral nervous and non-nervous system receptors, CNS and peripheral nervous and non-nervous system peptide transmitters, adhesion molecules, defensines, growth factors, vasoactive peptides and receptors, and binding proteins; or those mRNA which correspond to an oncogene. The present agent contains less than or about 15% adenosine (A), and in many cases is completely devoid of adenosine, and is also presented as a composition, which may be in the form of a capsule or cartridge, various formulations, and a kit provided with a delivery device and instructions for its use. The anti-sense-oligos of the invention may have their adenosine content reduced by substitution with an adenosine-like binding substitute such as universal base.

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The kit may also have other therapeutic agents, and ingredients for the composition. The agent also may be provided operatively linked to a vector and/or in a transfected host cell.

The agent, composition and formulation of the invention may be applied to the treatment of a disease or condition associated with the presence of an mRNA corresponding to at least one target gene, genomic flanking regions or proteins, by administration to a subject afflicted with the disease or condition of an amount of the MTA oligo of this invention effective to reduce the production or availability, or to increase the degradation, by the subject of at least one of the target mRNA. In a preferred embodiment, the agent may be administered in an amount effective to reduce the production or availability, or to increase the degradation of at least two of the target mRNAs. Typical diseases and conditions of this invention are those associated with impaired respiration and inflammation, including lung diseases, ailments and conditions that have a negative effect on the lungs of a subject. Examples of diseases and conditions, which may be treated preventively, prophylactically and therapeutically with the agent of this invention, are pulmonary vasoconstriction, inflammation, allergic rhinitis, acute asthma, allergies, asthma, impeded respiration, respiratory distress syndrome, pain, cystic fibrosis, pulmonary hypertension, pulmonary vasoconstriction, emphysema, chronic obstructive pulmonary disease (COPD), and cancers such as leukemias, lymphomas, carcinomas, and the like, e.g. colon cancer, breast cancer, lung cancer, pancreatic cancer, hepatocellular carcinoma, kidney cancer, melanoma, hepatic metastases, etc., as well as all types of cancers which may metastasize or have metastasized to the lung(s), including breast and prostate cancer. The present agent(s) is (are) also suitable for administration before, during and after other treatments, including radiation, chemotherapy, antibody therapy, phototherapy and cancer, and other types of surgery. Alternatively, the present agent may be effectively administered preventively, prophylactically or therapeutically, and in conjunction with other therapies, or by itself for conditions without known therapies or as a substitute for therapies that have significant negative side effects.

The composition of this invention may be administered by transdermal or systemic routes, including by, but not exclusively, oral, intracavitary, intranasal, intraanal, intravaginal, transdermal, intradermal, intrabuccal, intravenous, subcutaneous, intramuscular, intratumor, intraglandular, by inhalation, intraarterial, intravascular in general, into the ear, intracranial, intrathecal, intraorgan including via a shunt to, for example, the liver or other organs, by implantation and intraocular administration to a human or any other animal, including vertebrates, such as mammals. The treatment of this invention may be prophylactic or therapeutic. In a preferred embodiment, the present agents are administered directly into the respiratory system of a subject, so that the agent has direct access to the lungs, in an amount effective to reduce or inhibit the effect in the lung of the targeted diseases or conditions.

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The agent of this invention may be produced by selecting two or more targets such as genes, genomic flanking regions, intron-exon borders, and the like, or the entire sequence of RNAs, including non-coding RNAs, and RNAs encoding proteins known to be associated with at least one disease or condition; obtaining RNAs selected from the group consisting of RNAs corresponding to the genes and genomic flanking regions, and RNAs encoding the target proteins; selecting a segment of a first RNA which is at least about 60%, preferably about 80%, and still more preferably about 90% and even about 100% homologous to a segment of at least a segment of a second RNA; and synthesizing one or more anti-sense oligonucleotide(s) to the one or more RNA segments. The target RNA includes every segment of precursor and spliced mRNAs and other RNA molecules, including the 5'- and 3'-ends, and the coding portions as well as overlapping segments juxtaposed over the coding and non-coding sequences. In a preferred form of the invention, the two or more targets may be located in the same molecule. For instance, in the case of a mRNA encoding a protein with multiple subunits, MTA oligos may be found in segments of the RNA which encode different protein subunits.

Also provided are MTA oligos which are produced by the method of the invention.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

This invention arose from a desire by the inventor to improve on his own prior discovery, and those of others, that anti-sense oligonucleotides may be utilized therapeutically in the treatment of diseases or conditions which have multiple contributing pathways. The inventor reasoned that he could improve on his prior success in attenuating or enhancing the effects of one specific pathway by designing anti-sense oligonucleotides directed to a specific target associated with a disease or condition. He, thus, set out to attempt a novel and unobvious strategy directed to multiple targets. In so doing, he overcame numerous obstacles, particularly the extensive searching and selection necessary to obtain targets as well as their locating the desired sequences, be it genomic DNA, RNAs or proteins involved in specific diseases. Thereafter, he exemplified the invention by application to some specific diseases or conditions, and provided various preferred embodiments and specifically designed multiple targeted antisense oligonucleotide (MTA oligo) sequences.

The multi-targeted anti-sense (MTA) oligonucleotides of this invention have the capacity to attenuate the expression of more than one target mRNA, or to enhance or attenuate the activity of one or more pathways. By means of example, the present method may be practiced by first identifying all possible desadenosine (desA) anti-sense sequences of about 7, about 10, about 12, about 15, about 18, about 21 to about 28, about 30, about 35, about 40, about 45, about 50, about 60 or more mononucleotides in a target mRNA. This may be attained by searching for segments that are 7 or more nucleotides long within a target sequence

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which are low in, or lack, thymidine (T), a nucleotide which is complementary to adenosine (A). This search typically results in about 10 to 30 such desT segments, i. e. naturally lacking thymidine, or segments with low T content, e. g. up to and including about 15%T, from which anti-sense oligonucleotides of varying lengths may be designed for a typical target mRNA of average length, i. e. about 1800 nucleotides long. Thereafter, the sense sequence for each strictly complementary desA anti-sense sequence obtained for a specific target may be then deduced. The thus deduced sense sequence may be then used to search for sequences of preferred secondary targets. Alternatively, one or more sequence data bases, e. g., GENBANK, and the like, may be searched for alternative secondary sequences. Thus, the targeting may be undertaken in several manners, one being the selection of specific targets associated with one or more related diseases. Alternatively, a primary target may be selected first, and an anti-sense oligonucleotide found, preferably a desA oligonucleotide and, then, secondary, tertiary or more targets searched for. In a typical search, either the list of preferred secondary targets or of a data base, multiple instances of homologous secondary targets of interest are identified. That is, the present technology is directed to finding the instances where there are natural homologies between primary, secondary, and greater sequences, and utilizing the finding for the therapeutic treatment of specific diseases or conditions associated with the target macromolecules from which the MTAs are obtained.

The present technology relies on the design of anti-sense oligos targeted to mRNAs associated with ailments involving lung airway pathology(ies), and on their modification to reduce the occurrence of undesirable side effects caused by their release of adenosine upon breakdown, while preserving their activity and efficacy for their intended purpose. In this manner, the inventor targets a specific gene to design one or more anti-sense oligonucleotide(s) (oligos) that selectively bind(s) to the corresponding mRNA, and then reduces, if necessary, their content of adenosine via substitution with universal base or an adenosine analog incapable of activating adenosine A₁, A_{2b} or A₃ receptors. Based on his prior experience in the field, the inventor reasoned that in addition to "downregulating" specific genes, he could increase the effect of the agent(s) administered by either selecting segments of RNA that are devoid, or have a low content, of thymidine (T) or, alternatively, substitute one or more adenosine(s) present in the designed oligonucleotide(s) with other nucleotide bases, so called universal bases, which bind to thymidine but lack the ability to activate adenosine receptors and otherwise exercise the constricting effect of adenosine in the lungs, etc. Given that adenosine (A) is a nucleotide base complementary to thymidine (T), when a T appears in the RNA, the anti-sense oligo will have an A at the same position. For consistency's sake, all RNAs and oligonucleotides are represented in this patent by a single strand in the 5' to 3' direction, when read from left to right, although their complementary sequence(s) is (are) also encompassed

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within the four corners of the invention. In addition, all nucleotide bases and amino acids are represented utilizing the recommendations of the IUPAC-IUB Biochemical Nomenclature Commission, or by the known 3-letter code (for amino acids).

The method of the present invention may be used to treat ailments associated with reduced airway function in a subject, whatever its cause. The adenosine content of the antisense agent(s) of the invention have a reduced A content to prevent its liberation upon in vivo degradation of the agent(s). Examples of airway diseases that may be treated by the method of the present invention include cystic fibrosis, asthma, pulmonary hypertension and vasoconstriction, chronic obstructive pulmonary disease (COPD), chronic bronchitis, respiratory distress syndrome, lung cancer and lung metastatic cancers and other airway diseases, including those with inflammatory response.

Anti-sense oligos to the adenosine A₁, A_{2a}, A_{2b}, and A₃ receptors, CCR3 (chemokine receptors), bradykinin 2B, CAM (vascular cell adhesion molecule), and eosinophil receptors, among others, have been shown to be effective in down-regulating the expression of their genes. Some of these act to alleviate the symptoms or reduce respiratory ailments and/or inflammation, for example, by "down regulation" of the adenosine A1, A2a, A2b, and/or A3 receptors and CCR3, bradykinin 2B, VCAM (vascular cell adhesion molecule) and eosinophil receptors. These agents are preferably administered directly into the respiratory system, e.g., by inhalation or other means, so that they may reach the lungs without widespread systemic dissemination. This permits the use of substantially lower doses of the agent of the invention as compared with those administered by the prior art, systemically or by other generalized routes and, consequently, reduce undesirable side effects resulting from the agent's widespread distribution in the body. The agent(s) of this invention has (have) been shown to reduce the amount of receptor protein expressed by the tissue. These agents, thus, rather than merely interacting with their targets, e.g. a receptor, lower the number of target proteins that other drugs may interact with. In this manner, the present agent(s) afford(s) extremely high efficacy with low toxicity.

The adenosine receptors discussed above are mere examples of the high power of the inventor's technology. In fact, a large number of genes may be targeted in a similar manner by the present agent(s), to reduce or down-regulate protein expression. By means of example, if the target disease or condition is one associated with impeded or reduced breathing, bronchoconstriction, chronic bronchitis, pulmonary bronchoconstriction and/or hypertension, chronic obstructive pulmonary disease (COPD), allergy, asthma, cystic fibrosis, respiratory distress syndrome, cancers, which either directly or by metastasis afflict the lung, the present method may be applied to a list of potential target mRNAs, which includes the targets listed in Table 1 below, among others.

Table 1: Pulmonary Disease or Condition (Asthma/Inflammation) Targets

	TADIC A. I dillionary	
	Tra Train Footor	Interleukin-8 Receptor (IL-8 R)
	NfkB Transcription Factor	Interleukin-4 Receptor (IL-4R)
-	Interleukin-5 Receptor (IL-5R)	Interleukin-1β (IL-1β)
5	Interleukin-3 Receptor (IL-3R) Interleukin-1β Receptor (IL-1βR)	Eotaxin
		Major Basic Protein
	Tryptase β2-adrenergic Receptor Kinase	Endothelin Receptor A
	Endothelin Receptor B	Preproendothelin
10	Bradykinin B2 Receptor (B2BR)	IgE (High Affinity Receptor)
10	Interleukin-1 (IL-1)	Interleukin 1 Receptor (IL-1 R)
	Interleukin-9 (IL-9)	Interleukin-9 Receptor (IL-9 R)
	Interleukin-11 (IL-11)	Interleukin-11 Receptor (IL-11 R)
		Cyclogyugenase (COX)
15	Intercellular Adhesion Molecule 1 (ICAM	1) Vascular Cellular Adhesion Molecule (VCAM)
13	Rantes	Entionicital Leukocyte Municities 1997
	Cyclooxygenase-2 (COX-2)	GM-CSF, Endothelin-1
	Monocyte Activating Factor	Neutrophil Chemotactic Factor
	Neutrophil Elastase	Defensin 1,2,3
20	Muscarinic Acetylcholine Receptors	Platelet Activating Factor
20	Tumor Necrosis Factor α	5-lipoxygenase
	Phosphodiesterase IV	Substance P
	Substance P Receptor	Histamine Receptor
	Chymase	CCR-1 CC Chemokine Receptor
25	Interleukin-2 (IL-2)	Interleukin-4 (IL-4)
	Interleukin-12 (IL-12)	Interleukin-5 (IL-5)
	Interleukin-6 (IL-6)	Interleukin-7 (IL-7)
	Interleukin-8 (IL-8)	Interleukin-12 Receptor (IL-12R)
	Interleukin-7 Receptor (IL-7R)	Interleukin-1 (IL-1)
30	Interleukin-14 Receptor (IL-14R)	Interleukin-14
50	CCR-2 CC Chemokine Receptor	CCR-3 CC Chemokine Receptor
	CCR-4 CC Chemokine Receptor	CCR-5 CC Chemokine Receptor
	Prostanoid Receptors	GATA-3 Transcription Factor
	Neutrophil Adherence Receptor	MAP Kinase
35	Interleukin-15 (IL-15)	Interleukin-15 Receptor (IL-15R)
	Interleukin-11 (IL-11)	Interleukin-11 Receptor (IL-11R)
	NFAT Transcription Factors	STAT 4

Table 1: Pulmonary Disease or Condition (Asthma/Inflammation) Targets (Cont'ed)

	MIP-1α	MCP-2
	MCP-3	MCP-4
5	Cyclophillin (A, B, etc.)	Phospholipase A2
)	Basic Fibroblast Growth Factor	Metalloproteinase
	CSBP/p38 MAP Kinase	Tryptase Receptor
		Interleukin-3 (IL-3)
	PDG2 Interleukin-10 (IL-10)	Cyclosporin A - Binding Protein
Λ	EVENE Dinding Protein	α4β1 Selectin
0	FK506-Binding Protein	α4β7 Selectin
	Fibronectin	LFA-1 (CD11a/CD18)
	cMad CAM-1	LFA-1 Selectin
	PECAM-1	PSGL-1
-	C3bi	P-Selectin
15	E-Selectin	L-Selectin
	CD-34	Mac-1 (CD11b/CD18)
	p150,95	VLA-4
	Fucosyl transferase	CD11b/CD18
	CD-18/CD11a	C5a
20	ICAM2 and ICAM3	CCR1, CCR2, CCR4, CCR5
	CCR3 (Eotaxin Receptor)	AP-1 Transcription Factor
	LTB-4	Cysteinyl Leukotriene Receptor
	Protein kinase C	IrR Kinase 1 & 2
25	Tachykinnen Receptors (tach R)	(e.g., Substance P, NK-1 & NK-3 Receptors)
25	Interleukin-2 Receptor (IL-2R)	c-mas
	STAT 6 NF-Interleukin-6 (NF-IL-6)	Interleukin-10 Receptor (IL-10R)
		Interleukin-2 Receptor (IL-2R)
	Interleukin-3 (IL-3) Interleukin-13 (IL-13)	Interleukin-12 Receptor (IL-12R)
20	Interleukin-14 (IL-14)	Interleukin-6 Receptor (IL-6R)
30	Interleukin-16 (IL-16)	Interleukin-13 Receptor (IL-13R)
		Interleukin-16 Receptor (IL-16R)
	Medullasin Adenosine A ₁ Receptor (A ₁ R)	Tryptase-I
	Adenosine A_{1} Receptor $(A_{1} R)$ Adenosine A_{2b} Receptor $(A_{2b} R)$	Adenosine A ₃ Receptor (A ₃ R)
25		
35	β Tryptase	IgE Receptor β Subunit (IgE \mathbb{R} β)
	Adenosine A ₂ , Receptor (A ₂ , R) Fc-epsilon receptor CD23 antigen	IgE Receptor α Subunit (IgE R α)
	IgE Receptor Fc Epsilon Receptor (IgE	RFc & R) Substance P Receptor
	ige Receptor to Epsilon Receptor (1822)	I IVUIASC-1
40	Histidine decarboxylase	Eosinophil Cationic Protein
40	Prostaglandin D Synthase	Fosinophil Peroxidase
	Eosinophil Derived Neurotoxin Endothelial Nitric Oxide Synthase	Endothelial Monocyte Activating Factor
	Manage Factor	Cathensin G
	Neutrophil Oxidase Factor Macrophage Inflammatory Protein-1-	Interleukin-8 Receptor a Subunit (IL-8 R a)
45	Macrophage initiatinatory Protein-1	Substance P
45	Alpha/Rantes Receptor Endothelin Receptor ET-B	Endothelin ETA Receptor

The oligos of this invention may be obtained by first selecting fragments of a target nucleic acid having at least 4 contiguous nucleic acids selected from the group consisting of G and C, and then obtaining a first oligonucleotide 4 to 60 nucleotides long which comprises the selected fragment and has a C and G nucleic acid content of up to and including about 15%. The latter step may be conducted by obtaining a second oligonucleotide 4 to 60 nucleotides long comprising a sequence which is anti-sense to the selected fragment, the second oligonucleotide having an adenosine base content of up to and including about 15%. This method may also comprise, when the selected fragment comprises at least one thymidine base,

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substituting an adenosine base in the corresponding nucleotide of the anti-sense fragment with a universal base selected from the group consisting of heteroaromatic bases which bind to a thymidine base but have antagonist activity and less than about 0.3 of the adenosine base agonist activity at the adenosine A1, A2b and A3 receptors, and heteroaromatic bases which have no activity or have an agonist activity at the adenosine A22 receptor. The analogue heteroaromatic bases may be selected from all pyrimidines and purines, which may be substituted by O, halo, NH2, SH, SO, SO2, SO3, COOH and branched and fused primary and secondary amino, alkyl, alkenyl, alkynyl, cycloalkyl, heterocycloalkyl, aryl, heteroaryl, alkoxy, alkenoxy, acyl, cycloacyl, arylacyl, alkynoxy, cycloalkoxy, aroyl, arylthio, arylsulfoxyl, halocycloalkyl, alkylcycloalkyl, alkenylcycloalkyl, alkynylcycloalkyl, haloaryl, alkylaryl, alkenylaryl, alkynylaryl, arylalkyl, arylalkenyl, arylalkynyl, arylcycloalkyl, which may be further substituted by O, halo, NH2, primary, secondary and tertiary amine, SH, SO, SO₂, SO₃, cycloalkyl, heterocycloalkyl and heteroaryl. The pyrimidines and purines may be substituted at all positions as is known in the art, but preferred are those which are substituted at positions 1, 2, 3, 4, 7 and/or 8. More preferred are pyrimidines and purines such as theophylline, caffeine, dyphylline, etophylline, acephylline piperazine, bamifylline, enprofylline and xantine having the chemical formula

wherein R^1 and R^2 are independently H, alkyl, alkenyl or alkynyl and R^3 is H, aryl, dicycloalkyl, dicycloalkynyl, cycloalkyl, cycloalkenyl, cycloalkynyl, O-cycloalkynyl, O-cycloalkynyl, NH₂-alkylamino-ketoxyalkyloxy-aryl, mono and dialkylaminoalkyl-N-alkylamino-SO₂ aryl, among others.

When no segments having the desired T content were found or where desirable segments contained T, the inventor proposed to reduce the adenosine content of the anti-sense oligos corresponding to the thymidines (T) present in the target RNA to less than about 15%, or fully eliminated A from the oligonucleotide sequence as a means for preventing their breakdown products from freeing adenosine into the lung tissue environment and, thereby, aggravating the subject's ailment and/or countering the beneficial effect of the administered agent.

By means of example, the NfkB transcription factor may be selected as a primary target and searched for desthymidine (desT) segments. When a number of desT segments are found,

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the anti-sense segments may be deduced, and perhaps about 20 or even more desA anti-sense sequences obtained. These anti-sense sequences represent, when possible, all desA anti-sense sequences found within the mRNA of this primary target, and may be utilized to start the search for homologous sequences within a preferred list of secondary targets such as the one shown in Table 1 above or Table 2 below, or within a sequence data base, such as GENBANK. For each of the about 20 original desA anti-sense sequences found for NFk transcription factor, typically about 10 to 30 homologous sequences may be found among the other members of the group shown in Table 1 above (secondary, tertiary, and the like targets). In some instances, the search produces a homology for the primary target with, not only secondary targets (homology between primary target and sequence from one other target mRNA), but with tertiary targets (homology between primary target and sequences from, e. g. three other target mRNA), as well. The latter case, however, is more rare. When this occurs, the anti-sense oligos found are said to be 100% homologous. More typically, however, the sequences found contain some non-fully homologous nucleotides within the secondary or tertiary or quaternary sequences. In many cases, this mismatch would suffice to render the anti-sense oligonucleotide less active or even inactive against the target(s). In some instances, the presence of even one non-homologous nucleotide may be sufficient to reduce the activity of an anti-sense oligonucleotide. When the so called "homologous" sequences obtained have mismatches, acceptable are up to about 40%, preferably no more than about 30%, more preferable no more than 20%, still more preferable no more than 10% mismatched nucleotides. In some instances the higher % mismatches are acceptable, and the oligos still are active since the nonhomologous nucleotide may be "fixed" or replaced with a "Universal" base that may base-pair with similar or equal affinity with two or more of the four nucleotide present in natural DNA: A, G, C, and T. This "fixing" step generates a further novel sequence, different from the one found in nature, that permits the anti-sense oligonucleotide to bind, preferably equally well, with the primary target, the secondary target, the tertiary target, etc.

As the NfkB transcription factor is selected as a target, its mRNA or DNA are searched for low thymidine (T) or desthymidine (desT) fragments. Only desT segments of the mRNA or DNA are selected which, in turn, will produce desA anti-sense as their complementary strand. When a number of RNA desT segments are found, the sequence of the anti-sense segments may be deduced. Typically, about 10 to 30 and even larger numbers of desA anti-sense sequences may be obtained. These anti-sense sequences may include some or all desA anti-sense oligonucleotide sequences corresponding to desT segments of the mRNA of the target, such as anyone of those shown in Table 1 above or Table 2 below. When this occurs, the anti-sense oligonucleotides found are said to be 100% A-free. For each of the original desA anti-sense oligonucleotide sequences corresponding to the target gene, e.g. the NFkB

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transcription factor, typically about 10 to 30 sequences may be found within the target gene or RNA which have a low content of thymidine (RNA). In accordance with this invention, the selected fragment sequences may also contain a small number of thymidine (RNA) nucleotides within the secondary or tertiary or quaternary sequences. In some cases, a large adenosine content may suffice to render the anti-sense oligonucleotide less active or even inactive against the target. In accordance with this invention, these so called "non-fully desA" sequences may preferably have a content of adenosine of less than about 15%, more preferably less than about 10%, and still more preferably less than 5%, and some even less than 2% adenosine. In some instances a higher content of adenosine is acceptable and the oligonucleotides are still active, particularly where the adenosine nucleotide may be "fixed" or replaced with a "Universal" base that may base-pair with similar or equal affinity to two or more of the four nucleotide present in natural DNA: A, G, C, and T. A universal base is defined in this patent as any compound, more commonly an adenosine analogue, having the capacity to hybridize to thymidine, preferably having substantially reduced, or substantially lacking, ability to bind adenosine receptors. Alternatively, adenosine analogs which do not activate adenosine receptors, such as the adenosine A₁, A_{2 b} and/or A₃ receptors, most preferably A₁ receptors, may be used. One example of a universal base is α -deoxyribofuranosol-(5-nitroindole), and an artisan will know how to select others. This "fixing" step generates a further novel sequence, different from the one found in nature, that permits the anti-sense oligonucleotide to bind, preferably equally well, with the target RNA. An example of a universal base is 2deoxyribosyl-(5-nitroindole). Other examples of universal bases are 3-nitropyrrole-2'deoxynucleoside, 5-nitro-indole, 2-deoxyribosyl-(5-nitroindole), 2-deoxyribofuranosyl-(5nitroindole), 2'-deoxyinosine, 2'-deoxynebularine, 6H, 8H-3,4-dihydropyrimido [4,5-c] oxazine-7-one and 2-amino-6-methoxyaminopurine. In addition to the above, Universal bases which may be substituted for any other base although with somewhat reduced hybridization potential, include 3-nitropyrrole 2'-deoxynucleoside 2-deoxyribofuranosyl-(5-nitroindole), 2'-deoxyinosine and 2'-deoxynebularine (Glen Research, Sterling, VA). More specific mismatch repairs may be made using "P" nucleotide, 6H, 8H-3, 4-dihydropyrimido[4,5-c] [1,2] oxazin-7-one, which base pairs with either guanine (G) or adenine (A) and "K" nucleotide, 2-amino-6-methoxyaminopurine, which base pairs with either cytidine (C) or thymidine (T), among others. Others which are known in the art are also suitable. See, for example, Loakes, D. and Brown, D. M., Nucl . Acids Res. 22:4039-4043 (1994); Ohtsuka, E. et al., J. Biol. Chem.260(5):2605-2608 (1985); Lin, P.K.T. and Brown, D. M., Nucleic Acids Res. 20(19):5149-5152 (1992; Nichols, R. et al., Nature 369(6480): 492-493 (1994); Rahmon, M. S. and Humayun, N. Z., Mutation Research 377 (2): 263-8 (1997); Amosova, O., et al., Nucleic Acids Res. 25 (!0): 1930-1934 (1997); Loakes D. & Brown, D. M., Nucleic

Acids Res. 22 (20): 4039-4043 (1994), the entire sections relating to universal bases and their preparation and use in nucleic acid binding is incoorporated herein by reference.

When non-fully desT sequences are found in the naturally occurring target, they typically are selected so that about 1 to 3 universal base substitutions will suffice to obtain a 100% "desA" anti-sense oligonucleotide. Thus, the present method provides either anti-sense oligonucleotides to different targets which are low in, or devoid of, A content, as well as anti-sense oligonucleotides where one or more adenosine nucleotides, e. g. about 1 to 3, or more, may be "fixed" by replacement with a "Universal" base. Universal bases are known in the art and need not be listed herein. An artisan will know which bases may act as universal bases, and replace them for A.

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The present approach to the design of anti-sense oligonucleotide approach is also applicable to a variety of other diseases or conditions, including other inflammatory diseases, such as cystic fibrosis, chronic obstructive pulmonary disease, chronic bronchitis, pulmonary hypertension, cancers, including those which metastasize to the lung, such as breast cancer, colon cancer, respiratory distress syndrome, prostate cancer, pancreatic cancer, kidney cancer, lymphomas, melanomas, hepatocellular carcinomas, etc.

As used herein, the term "treat" or "treating" asthma or other respiratory and inflammatory conditions or diseases refers to a treatment which decreases the likelihood that the subject administered such treatment will manifest symptoms of a respiratory or inflammatory lung disease or other lung conditions. The term "down-regulate" refers to inducing a decrease in production, secretion or availability (and thus a decrease in concentration) of the targeted intracellular protein.

The present invention is concerned primarily with the treatment of vertebrates, and within this group, of mammals, including human and non-human simians, wild and domesticated animals, marine and land animals, household pets, and zoo animals, for example, felines, canines, equines, pachiderms, cetaceans, and still more preferably to human subjects. One particularly suitable application of this technology is for veterinary purposes, and includes all types of small and large animals in the care of a veterinarian, including wild animals, marine animals, household animals, zoo animals, and the like. Targeted genes and proteins are preferably mammalian, and the sequences targeted are preferably of the same species as the subject being treated. Although in many instances, targets of a different species are also suitable, particularly those segments of the target RNA or gene that display greater than about 45% homology, preferably greater than about 85% homology, still more preferably greater than about 95% homology, with the recipient's sequence. A preferable group of agents is composed of des-A anti-sense oligos. Another preferred group is composed of non-fully desA oligonucleotides, where one or more adenosine bases are replaced with universal bases.

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The terms "anti-sense" oligonucleotides generally refers to small, synthetic oligonucleotides, resembling single-stranded DNA, which in this patent are applied to the inhibition of gene expression by inhibition of a target messenger RNA (mRNA). See, Milligan, J. F. et al., J. Med. Chem. 36(14), 1923-1937 (1993), the relevant portion of which is hereby incorporated in its entirety by reference. The present agents inhibit gene expression of target genes, such as those of the adenosine A_1 A_{2a} , A_{2b} , or A_3 receptors, CCR3 (chemical receptor 320, also known as the eotaxin receptor), VCAM (vascular cell adhesion molecule), eonophil receptor, bradykinin 2B receptor, and many others listed in Table 1 above. This is generally attained by hybridization of the anti-sense oligonucleotides to coding (sense) sequences of a targeted messenger RNA (mRNA), as is known in the art. The exogenously administered agents of the invention decrease the levels of mRNA and protein encoded by the target gene and/or cause changes in the growth characteristics or shapes of the thus treated cells. See, Milligan et al. (1993); Helene, C. and Toulme, J. Biochim. Biophys. Acta 1049, 99-125 (1990); Cohen, J. S. D., Ed., Oligodeoxynucleotides as Anti-sense Inhibitors of Gene Expression; CRC Press: Boca Raton, FL (1987), the relevant portion of which is hereby incorporated in its entirety by reference. As used herein, "anti-sense oligonucleotide" is generally a short sequence of synthetic nucleotide that (1) hybridizes to any segment of a mRNA encoding a targeted protein under appropriate hybridization conditions, and which (2) upon hybridization causes a decrease in gene expression of the targeted protein.

The terms "des-adenosine" (desA) and "des-thymidine" (desT) refer to oligonucleotides substantially lacking either adenosine (desA) or thymidine (desT). In some instances, the des T sequences are naturally occurring, and in others they may result from substitution of an undesirable nucleotide (A) by another one lacking its undesirable activity. In the present context, the substitution is generally accomplished by substitution of A with a "universal base", as is known in the art.

The mRNA sequence of the targeted protein may be derived from the nucleotide sequence of the gene expressing the protein. For example, the sequence of the genomic human adenosine A₁ receptor and that of the rat and human adenosine A₂ receptors are known. See, US Pat. No. 5,320,962; Zhou, F., et al., Proc. Nat'1 Acad. Sci. (USA) 89:7432 (1992); Jacobson, M.A., et al., U.K. Pat. Appl. No. 9304582.1. The sequence of the adenosine A_{2b} receptor gene is also known. See, Salvatore, C. A., Luneau, C. J., Johnson, R. G. and Jacobson, M., Genomics (1995), the relevant portion of which is hereby incorporated in its entirety by reference. The sequences of many of the exemplary target genes are also known. See, GenBank, NIH. The sequences of those genes whose sequences are not yet available may be obtained by isolating the target segments applying technology known in the art. Once the sequence of the gene, its RNA and/or the protein are known, an anti-sense oligonucleotides

may be produced according to this invention as described above to reduce the production of the targeted protein in accordance with standard techniques.

In one aspect of this invention, the anti-sense oligonucleotide has a sequence which specifically binds to a portion or segment of an mRNA molecule which encodes a protein associated with a disease or condition associated with impeded breathing, lung inflammation, airway obstruction, bronchitis, and the like. One effect of this binding is to reduce or even prevent the translation of the corresponding mRNA and, thereby, reduce the available amount of target protein in the subject's lung.

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In one preferred embodiment of this invention, the phosphodiester residues of the antisense oligonucleotide are modified or substituted. Chemical analogs of oligonucleotides with modified or substituted phosphodiester residues, e.g., to the methylphosphonate, the phosphotriester, the phosphorothioate, the phosphorodithioate, or the phosphoramidate, which increase the in vivo stability of the oligonucleotide are particularly preferred. The naturally occurring phosphodiester linkages of oligonucleotides are susceptible to some degree of degradation by cellular nucleases. Many of the residues proposed herein, on the contrary, are highly resistant to nuclease degradation. See Milligan et al., and Cohen, J. S. D., supra. In another preferred embodiment of the invention, the oligonucleotides may be protected from degradation by adding a " 3'-end cap" by which nuclease-resistant linkages are substituted for phosphodiester linkages at the 3' end of the oligonucleotide. See, Tidd, D. M. and Warenius, H.M., Be. J. Cancer 60: 343-350 (1989); Shaw, J.P. et al., Nucleic Acids Res. 19: 747-750 (1991), the relevant section of which are incorporated in their entireties herein by reference. Phosphoramidates, phosphorothioates, and methylphosphonate linkages all function adequately in this manner for the purposes of this invention. The more extensive the modification of the phosphodiester backbone the more stable the resulting agent, and in many instances the higher their RNA affinity and cellular permeation. See Milligan, et al., supra. Thus, the number of residues which may be modified or substituted will vary depending on the need, target, and route of administration, and may be from 1 to all the residues, to any number in between. Many different methods for replacing the entire phosphodiester backbone with novel linkages are known. See, Millikan et al, supra. Preferred backbone analogue residues include phosphorothioate, methylphosphonate, phosphotriester, thioformacetal, phosphorodithioate, phosphoramidate, formacetal boranophosphate, 3'-thioformacetal, 5'-thioether, carbonate, 5'-N-carbamate, sulfate, sulfonate, sulfonamide, sulfone, sulfite., 2'-O methyl, methylene(methylimino) (MMI). sulfide, hydroxylamine, sulfoxide, methyleneoxy(methylimino) (MOMI) residues. Phosphorothioate and methylphosphonatemodified oligonucleotides are particularly preferred due to their availability through automated oligonucleotide synthesis. See, Millikan et al, supra. Where appropriate, the agent of this

invention may be administered in the form of their pharmaceutically acceptable salts, or as a mixture of the anti-sense oligonucleotide and its salt. In another embodiment of this invention, a mixture of different anti-sense oligonucleotides or their pharmaceutically acceptable slats is administered.

The agents of this invention have the capacity to attenuate the expression of one target mRNA and/or to enhance or attenuate the activity of one pathway. By means of example, the present method may be practiced by identifying all possible deoxyribonucleotide segments which are low in thymidine (T) or deoxynucleotide segments low in adenosine (A) of about 7 or more mononucleotides, preferably up to about 60 mononucleotides, more preferably about 10 to about 36 mononucleotides, and still more preferably about 12 to about 21 mononucleotides, in a target mRNA or a gene, respectively. This may be attained by searching for mononucleotide segments within a target sequence which are low in, or lack thymidine (RNA), a nucleotide which is complementary to adenosine, or that are low in adenosine (gene), that are 7 or more nucleotides long. In most cases, this search typically results in about 10 to 30 such sequences, I. e. naturally lacking or having less than about 40% adenosine, anti-sense oligonucleotides of varying lengths for a typical target mRNA of average length, i. e., about 1800 nucleotides long. Those with high content of T or A, respectively, may be fixed by substitution of a universal base for one or more As.

The agent(s) of this invention may be of any suitable length, including but not limited to, about 7 to about 60 nucleotides long, preferably about 12 to about 45, more preferably up to about 30 nucleotides long, and still more preferably up to about 21, although they may be of other lengths as well, depending on the particular target and the mode of delivery. The agent(s) of the invention may be directed to any and all segments of a target RNA. One preferred group of agent(s) includes those directed to an mRNA region containing a junction between an intron and an exon. Where the agent is directed to an intron/exon junction, it may either entirely overlie the junction or it may be sufficiently close to the junction to inhibit the splicing-out of the intervening exon during processing of precursor mRNA to mature mRNA, e.g. with the 3' or 5' terminus of the anti-sense oligonucleotide being positioned within about, for example, within about 2 to 10, preferably about 3 to 5, nucleotide of the intron/exon junction. Also preferred are anti-sense oligonucleotides which overlap the initiation codon, and those near the 5' and 3' termini of the coding region.

This multi-targeted anti-sense (MTA) oligonucleotide approach is, thus, applicable to a variety of other diseases or conditions, including other inflammatory diseases, such as cystic fibrosis, chronic obstructive pulmonary disease, chronic bronchitis, etc.. Other specific diseases or conditions to which this technology is effectively applied are pulmonary hypertension, and cancers.

Table 2 below provides a number of targets to which multi-targeted anti-sense (MTA) oligonucleotides are effectively applied. Others may also be targeted.

Table 2: Cancer Targets

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Transforming Oncogenes	Therapy Targets
ras	thymidylate synthetase
src	thymidylate synthetase
myc	dihydrofolate reductase
bcl-2	thymidine kinase
	deoxycytidine kinase
	ribonucleotide reductase

A group of preferred targets for the treatment of cancers are genes associated with different types of cancers, or those generally known to be associated with malignancies, whether they are regulatory or involved in the production of RNA and/or proteins. Examples are transforming oncogenes, including, but not limited to, ras, src, myc, and bcl-2, among others. Other targets are those to which present cancer chemotherapeutic agents are directed to, such as various enzymes, primarily, although not exclusively, thymidylate synthetase, dihydrofolate reductase, thymidine kinase, deoxycytidine kinase, ribonucleotide reductase, and the like.

The present technology is extremely important for the treatment of diseases or conditions such as cancer given that traditional cancer therapies are fraught with the unresolved problem of selectively killing cancer cells while preserving normal living cells from the devastating effects of treatments such as chemotherapy, radiotherapy, and the like. The present technology provides the ability of simultaneously attenuating or enhancing multiple pathways. This approach provides a significant advantage for the treatment of cancer because it permits the selection of a combination of multiple pathways, including primary, secondary and possibly tertiary targets, which are not generally expressed simultaneously in normal cells. Thus, the present agent may be administered to a subject to cause a selective increase in toxicity within tumor cells that, for instance, express all three targets while normal cells that may expresses only one or two of the targets will be significantly less affected or even spared.

This invention thus provides an agent, comprising an anti-sense oligonucleotide to two or more mRNAs selected from the group consisting of RNAs corresponding to target genes, to genomic flanking regions, the initiation origin, intron-exon borders, and the like, or the entire sequence of precursor RNAs, including the coding region of mRNAs, non-coding RNA

segments, the 5'-end and the 3'-end, e.g. poly-A segment and oligos targeted to the juxtasection between coding and non-coding regions, and RNAs encoding proteins known to be associated with one or more diseases or conditions or mixtures thereof.

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The agents administered in accordance with this invention are preferably designed to be anti-sense to target genes and/or mRNAs related in origin to the species to which it is to be administered. When treating humans, the agents are preferably designed to be anti-sense to a human gene or RNA. The agents of the invention encompass oligonucleotides which are antisense to naturally occurring DNA and/or RNA sequences, fragments thereof of up to a length of one (1) base less than the targeted sequence, preferably at least about 7 nucleotides long, oligos having only over about 0.02%, more preferably over about 0.1%, still more preferably over about 1%, and even more preferably over about 4% adenosine nucleotides, and up to about 30%, more preferably up to about 15%, still more preferably up to about 10% and even more preferably up to about 5%, adenosine nucleotide, or lacking adenosine altogether, and oligos in which one or more of the adenosine nucleotides have been replaced with so-called universal bases, which may pair up with thymidine nucleotides but fail to substantially trigger adenosine receptor activity. Examples of human sequences and fragments, which are not limiting, of anti-sense oligonucleotide of the invention are the following fragments as well as shorter segments of the fragments and of the full gene or mRNA coding sequences, exons and intron-exon junctions encompassing preferably 7, 10, 15, 18 to 21, 24, 27, 30, n-1 nucleotides for each sequence, where n is the sequence's total number of nucleotides. These fragments may be selected from any portion of the longer oligo, for example, from the middle, 5'- end, 3'- end or starting at any other site of the original sequence. Of particular importance are fragments of low adenosine nucleotide content, that is, those fragments containing less than or about 30%, preferably less than or about 15%, more preferably less than or about 10%, and even more preferably less than or about 5%, and most preferably those devoid of adenosine nucleotide, either by choice or by replacement with a universal base in accordance with this invention. The agent of the invention includes as a most preferred group sequences and their fragments where one or more adenosines present in the sequence have been replaced by a universal base (B), as exemplified here. Similarly, also encompassed are all shorter fragments of the B-containing fragments designed by substitution of B(s) for adenosine(s) (A(s)) contained in the sequences, fragments thereof or segments thereof, as described above. A limited list of sequences and fragments is provided below.

Some of the examples of anti-sense oligonucleotide sequence fragments target the initiation codon of the respective gene, and in some cases adenosine is substituted with a universal base adenosine analogue denoted as "B", which lacks ability to bind to the adenosine A_1 and/or A_3 receptors. In fact, such replacement nucleotide acts as a "spacer". Many of the

examples shown below provide one such sequence and many fragments overlapping the initiation codon, preferably wherein the number of nucleotides n is about 7, about 10, about 12, about 15, about 18, about 21 and up to about 28, about 35, about 40, about 50, about 60.

In one embodiment, at least one of the mRNAs to which the MTA oligo of the invention is targeted encodes a protein such as transcription factors, stimulating and activating factors, intracellular and extracellular receptors and peptide transmitters in general, interleukins, interleukin receptors, chemokines, chemokine receptors, endogenously produced specific and non-specific enzymes, immunoglobulins, antibody receptors, central nervous system (CNS) and peripheral nervous and non-nervous system receptors, CNS and peripheral nervous and non-nervous system peptide transmitters, adhesion molecules, defensines, growth factors, vasoactive peptides and receptors, and binding proteins, among others; or the mRNA is corresponding to an oncogene and other genes associated with various diseases or conditions.

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Examples of target proteins are eotaxin, major basic protein, preproendothelin, eosinophil cationic protein, P-selectin, STAT 4, MIP-1α, MCP-2, MCP-3, MCP-4, STAT 6, c-mas, NF-IL-6, cyclophillins, PDG2, cyclosporin A-binding protein, FK5-binding protein, fibronectin, LFA-1 (CD11a/CD18), PECAM-1, C3bi, PSGL-1, CD-34, substance P, p150,95, Mac-1 (CD11b/CD18), VLA-4, CD-18/CD11a, CD11b/CD18, C5a, CCR1, CCR2, CCR4, CCR5, and LTB-4, among others. Others are, however, suitable, as well.

In another embodiment, at least one of the mRNAs to which the MTA oligo is targeted encodes intracellular and extracellular receptors and peptide transmitters such as sympathomimetic receptors, parasympathetic receptors, GABA receptors, adenosine receptors, bradykinin receptors, insulin receptors, glucagon receptors, prostaglandin receptors, thyroid receptors, androgen receptors, anabolic receptors, estrogen receptors, progesterone receptors, receptors associated with the coagulation cascade, adenohypophyseal receptors, adenohypophyseal peptide transmitters, and histamine receptors (HisR), among others. However others are also contemplated.

The encoded sympathomimetic receptors and parasympathomimetic receptors include acetylcholinesterase receptors (AcChaseR) acetylcholine receptors (AcChR), atropine receptors, muscarinic receptors, epinephrine receptors (EpiR), dopamine receptors (DOPAR), and norepinephrine receptors (NEpiR), among others. Further examples of encoded receptors are adenosine A_1 receptor, adenosine A_2 B receptor, adenosine A_3 receptor, endothelin receptor A, endothelin receptor B, IgE high affinity receptor, muscarinic acetylcholine receptors, substance P receptor, histamine receptor, CCR-1 CC chemokine receptor, CCR-2 CC chemokine receptor, CCR-3 CC chemokine receptor (Eotaxin Receptor), interleukin-1 β receptor (IL-1 β R), interleukin-1 receptor (IL-1R), interleukin-1 β receptor (IL-1 β R), interleukin-3 receptor (IL-3R), CCR-4 CC chemokine receptor, cysteinyl leukotriene

receptors, prostanoid receptors, GATA-3 transcription factor receptor, interleukin-1 receptor (IL-1R), interleukin-4 receptor (IL-4R), interleukin-5 receptor (IL-5R), interleukin-8 receptor (IL-8R), interleukin-9 receptor (IL-9R), interleukin-11 receptor (IL-11R), bradykinin B2 receptor, sympathomimetic receptors, parasympathomimetic receptors, GABA receptors, adenosine receptors, bradykinin receptors, insulin receptors, glucagon receptors, prostaglandin receptors, thyroid receptors, androgen receptors, anabolic receptors, estrogen receptors, progesterone receptors, receptors associated with the coagulation cascade, adenohypophyseal receptors, and histamine receptors (HisR). Others are also contemplated even though not listed herein.

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The encoded enzymes for development of the MTA oligos of the invention include synthetases, kinases, oxidases, phosphatases, reductases, polysaccharide, triglyceride, and protein hydrolases, esterases, elastases, and, polysaccharide, triglyceride, lipid, and protein synthases, among others. Examples of target enzymes are tryptase, inducible nitric oxide synthase, cyclooxygenase (Cox), MAP kinase, eosinophil peroxidase, β2-adrenergic receptor kinase, leukotriene c-4 synthase, 5-lipooxygenase, phosphodiesterase IV, metalloproteinase, tryptase, CSBP/p38 MAP kinase, neutrophil elastase, phospholipase A2, cyclooxygenase 2 (Cox-2), fucosyl transferase, chymase, protein kinase C, thymidylate synthetase, dihydrofolate reductase, thymidine kinase, deoxycytidine kinase, and ribonucleotide reductase, among others. Any enzyme associated with a disease or condition, however, is suitable as a target for this invention.

Suitable encoded factors for application of this invention are, among others, NfkB transcription factor, granulocyte macrophage colony stimulating factor (GM-CSF), AP-1 transcription factor, GATA-3 transcription factor, monocyte activating factor, neutrophil chemotactic factor, granulocyte/macrophage colony-stimulating-factor (G-CSF), NFAT transcription factors, platelet activating factor, tumor necrosis factor α (TNF α), and basic fibroblast growth factor (BFGF). Additional factors are also within the invention even though not specifically mentioned.

Suitable adhesion molecules for use with this invention include intracellular adhesion molecules 1 (ICAM-1), 2 (ICAM-2) and 3 (ICAM-3), vascular cellular adhesion molecule (VCAM), endothelial leukocyte adhesion molecule-1 (ELAM-1), neutrophil adherence receptor, mad CAM-1, and the like. Other known and unknown factors (at this time) may also be targeted herein.

Among the cytokines, lymphokines and chemokines preferred are interleukin-1 (IL-1), interleukin-1β (IL-1)interleukin-3 (IL-3), interleukin-4 (IL-4), interleukin-5 (IL-5), interleukin-8 (IL-8), interleukin-9 (IL-9), interleukin-11 (IL-11), CCR-5 CC chemokine, and Rantes. Others, however, may also be targeted, as they are known to be involved in specific diseases

or conditions to be treated, or for their generic activities, such as inflammation.

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Examples of defensins for the practice of this invention are defensin 1, defensin 2, and defensin 3, and of selectins are $\alpha 4\beta 1$ selectin, $\alpha 4\beta 7$ selectin, LFA-1 selectin, E-selectin, P-selectin, and L-selectin. Examples of oncogenes, although not an all inclusive list, are ras, src, myc, and bcl-2. Others, however, are also suitable for use with this invention.

The agents administered in accordance with this invention are preferably designed to be anti-sense to target genes and/or mRNAs related in origin to the species to which it is to be administered. When treating humans, the agents are preferably designed to be anti-sense to a human gene or RNA. The agents of the invention encompass oligonucleotides which are antisense to naturally occurring DNA and/or RNA sequences, fragments thereof of up to a length of one (1) base less than the targeted sequence, preferably at least about 7 nucleotides long, oligos having only over about 0.02%, more preferably over about 0.1%, still more preferably over about 1%, and even more preferably over about 4% adenosine nucleotides, and up to about 30%, more preferably up to about 15%, still more preferably up to about 10% and even more preferably up to about 5%, adenosine nucleotide, or lacking adenosine altogether, and oligos in which one or more of the adenosine nucleotides have been replaced with so-called universal bases, which may pair up with thymidine nucleotides but fail to substantially trigger adenosine receptor activity. Examples of human sequences and fragments, which are not limiting, of anti-sense oligonucleotide of the invention are the following fragments as well as shorter segments of the fragments and of the full gene or mRNA coding sequences, exons and intron-exon junctions encompassing preferably 7, 10, 15, 18 to 21, 24, 27, 30, n-1 nucleotides for each sequence, where n is the sequence's total number of nucleotides. These fragments may be selected from any portion of the longer oligo, for example, from the middle, 5'- end, 3'- end or starting at any other site of the original sequence. Of particular importance are fragments of low adenosine nucleotide content, that is, those fragments containing less than or about 30%, preferably less than or about 15%, more preferably less than or about 10%, and even more preferably less than or about 5%, and most preferably those devoid of adenosine nucleotide, either by choice or by replacement with a universal base in accordance with this invention. The agent of the invention includes as a most preferred group sequences and their fragments where one or more adenosines present in the sequence have been replaced by a universal base (B), as exemplified here. Similarly, also encompassed are all shorter fragments of the B-containing fragments designed by substitution of B(s) for adenosine(s) (A(s)) contained in the sequences, fragments thereof or segments thereof, as described above. A limited list of sequences and fragments is provided below.

Some of the examples of anti-sense oligonucleotide sequence fragments target the initiation codon of the respective gene, and in some cases adenosine is substituted with a

universal base adenosine analogue denoted as "B", which lacks ability to bind to the adenosine A₁ and/or A₃ receptors. In fact, such replacement nucleotide acts as a "spacer". Many of the examples shown below provide one such sequence and many fragments overlapping the initiation codon, preferably wherein the number of nucleotides n is about 7, about 10, about 12, about 15, about 18, about 21 and up to about 28, about 35, about 40, about 50, about 60.

Human Receptor-related Antisense Polynucleotide

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5'--GGCGGCCTGG AAAGCTGAGA TGGAGGGCGG CATGGCGGGC ACAGGCTGGG C TGCTTTCT TTTCTGGGCC GTGGTTCTTG CCCTCCTTTG GCTGCCGTGC CCGCTCCCCG GCCTCCTGGC GGGTGGCCGT TGGGCCCGTG TTCCCCTGGG GCCTGGGGCT CCCTTCTCTC GCCCTTCTTG CTGGGCCTCT GCTGCTGCTG GTGCTGTGGC CCCCGTACA CCGAGGAGCC CATGATGGGC ATGCCACAGA CGACAGGCGT BCBCCGBGGB GCCCBTGBTG GGCBTGCCBC BGBCGBCBGG C GGC GCC GTG CCG CTT CTG GCT GGG CCC CGG GCG CCC CCT CCC CTC TTG CTC GGG TCC CCG TG ACA GCG CGT CCT GTG TCT CCA GCA GCA TGG CCG GGC CAG CTG GGC CCC BCB GCG CGT CCT GTG TCT CCB GCB GCB TGG CCG GGC CBG CTG GGC CCC ACA GAG CAG TGC TGT TGT TGG GCA TCT TGC CTT CCC AGG G BCB GBG CB TGC TGT TGT TGG GCB TCT TGC CTT GGT CIT CC CTC CTG CTC TTT TTT C ATT TGC TCT CCT ATT ACT TTC TGT GTC CAT TTT TTC ATT AAC CGA GCT GT BTT TGC TCT CCT BTT BCT TTC TGT GTC CBT TTT TTC BTT BBC CGB GCT GT GCC TGT GTC TGT CCT CCT GCT TCG TTC TCT TO GCT GCC TCC GTT TGG GTG GC TCT CTG AAT ATT GAC CTT CCT CCA TGG CGG TCC TGC TTG GAT TCT CCC GA TET CTG BBT BTT GBC CTT CCT CCB TGG CGG TCC TGC TTG GBT TCT CCC GB GCC TTT CCT GGT TCT CTT GTT GTT TTT GGG GTT TGG CTT ACA GTA GAG TAG GGG ATT CCA TGG CAG GAG CCA TCT TCT TCA TGG ACT CC TTC AAG GAG ACC TTA GGT TTC TGA GGG ACT GCT AAC ACG CCA TCT GGA GC BCB GTB GBG TBG GGG BTT CCB TGG CBG GBG CCB TCT TCT TCB TGG BCT CC TTC BBG GBG BCC TTB GGT TTC TGB GGG BCT GCT BBC BCG CCB TCT GGB GC GTT GTT TIT GGG GTT TGG CTT GCC TIT CCT GGT TCT CTT BCB GTB GBG TBG GGG BTT CCB TGG CBG GBG CCB TCT TCT TCB TGG BCT CC TTC BBG GBG BCC TTB GGT TTC TGB GGG BCT GCT BBC BCG CCB TCT GGB GC GCC TGT GTC TGT CCT CCT GCT TCG TTC CTC TCG CTT GGT GCC CTT GCC G GTC CTG CTC CGG GCT GTG G GTC CTC GCC CTG 30 CGT GGT TGG GGG TCT TC GCT GCC TCC GTT TGG GTG GC GAT CTC TGA ATA TTGA CCT TCC ATG GCG GTC CTG CTT GGA GBT CTC TGB BTB TTGB CCT TCC BTG GCG GTC CTG CTT GGB TCT GGG GTG TCC TGG CCT TCG TGG TTC CTC TTC CIT CGT TTG CCG TCC GCG GGG GCC CCC GGG CCT GGC TGC GCT CCT GCC CCG CCT CTT TCC CGG GCT CTT GCG CTG 35 CGT GGT TCC TCT TCC TTC GTT TGC CGT CCG CGG GGG CCC CCG GGC CT GGC TGC GCT CCT GCC CCG CCT CTT TCC GAC AGG GCA GGG CGA TCA GGA GCA GCG TGA GCC AAA GGA GGA CCA TCG GGA ACG CAG CTC CGG AAC GCA GGA 40 CAG AGG TGC C GC BGG BGB CBG GGC BGG GCG BTC BGG BGC BGC GTG BGC CBB BGG BGG BCC BTC GGG BBC GCB GCT CCG GBB CGC BGG BCB GBG GTG CC TCT GCC CTG TCC GCC GGC TCT TCG GTG GCT CGG CCC CGC TCC TTG TCT TGC CGC GGG TTG GTT CCT GGG CCT GGT TCT TGC GGG CGT TTC GGT CTG GCT GGT CTG GGC CCG CGG TGC GGC 45 TGG CTG CTT CGG GCC GCC TGG GCT TCC CTG TGC CCC TTT CCT CTG GGT CCC CCT CCC GTT CCA AGC TGC ACC GCA CAG ACC GGC GCT ACA GGA CAG AGC CAG GCA AGC ACC CAT GGG GAT CCA GGC CCA GCT GTT CCB BGC TGC BCC GCB CBG BCC GGC GCT BCB GGB CBG BGC CBG GCB BGC CCB GCC CBT GGG GBT CCB GGC CCB GCT G CTCAGTGGCC CCCAAAAGGA TGAGTAATAC ATGCGCCACG ATGATCATAT CCTTTTTACT ATGAGGCCGT GTCTGTCGTG 50

GCCTCTCTGC -GGGGGTGGCT TCCTGCCGCG TCTCTGGGCC GTCCCGTCCC TCGGCCCCGC GCCGCGCTCG GCTCCTCTCC GGCTGCGCGC TGCTGGCTGC CCTGCTGGCC GCGCCGGGGC CTGTCCGCCT CTGCGGGGCGC TGTCTCCTGG CTTGTCTTCC 5 CCTCCGGGGC GGGGGTTTCT GGCCGTGGGG GTCTTGCCTG GCCTCCGGGC TCCTGCTTGT CTTGCCTTCC TTCTCTGGTC GGTTGTGGCT CGGGGCTCCG TGGGTCCCTG GCGCCCGTTT GTGTTTTGTC TTTTCCCCTG GCGTCCCTGT GCCCCTCTCC TETECTTECT CTGCTTCTCG CTCTCCTTTG TGGGGCCCTC CCTGCTGCTC TTGGTTTTGG GCTTTTTTTC TCTTCCTCCT TTTTCGTGCG TGGGCCTCC GCACGCCTCT TGCCACCTCC TGCGCAGGGC AGCGCCTTGG GGCCAGCGCC GCTCCCGGCG 10 CGGCCAGCAG GGCAGCCAGC AGCGCGCAGC CGACGGCCAG CATGCTTCCT CCTCGGCTAC CACTCCATGG TCCCGCAGAG GCGGACAGGC GCBCGCCTC TTGCCBCCTC CTGCGCBGGG CBGCGCCTTG GGGCCBGCGC CGCTCCCGGC GCGGCCBGCB GGGCBGCCBG CBGCGCCBG CCGBCGGCCB GCBTGCTTCC TCCTCGGCTB CCBCTCCBTG GTCCCGCBGB GGCGGBCBGG C GCTGCCCGGC GGGGTGTGCG CTTGGCGCTC CCGTGCTCGG TTCTCTGTCT CCCGGTCCCC CTTGCCTGGC GTCTCGGGCC TTCGTCCTCT TCCTCTTCTT CCTTCCGCTC CGTGGGGGCT GCTTGGTGGG GGCCTGTGCCT CGGGGTCCCG GGGCTTCTGG 15 CCCTTGCCGT TCATGGTGGC TAGGTGGGGC GTTCBTGGTG GCTBGGTGGG GC GGG GTG GGT BGG CCG TGT CTG GGGGTT 20 GGT TGC CGGG CCC GCG GCT GCA GGG G ACAGGGGCTG TAATCTTCATC TGCAGGTGGC ATGCCAGTGA AATTTAGATC ATCAAAATCC CACATCTGTG GATCTGTAAT ATTTGACATG TCCTCTTCAG TTTCAGCAAT GGTTTGATCT AACTGAAGCA CCGGCCAGGB CBGGGGCTGT BBTCTTCBTC TGCBGGTGGC BTGCCBGTGB BBTTTBGBTC BTCBBBBTCC CBCBTCTGTG GBTCTGTBBT BTTTGBCBTG TCCTCTTCBG TTTCBGCBB TGGTTTGBTC TBBCTGBBGC BCCGGCCBGG TGGCTCGGTG CTTCTGCCCC TGTTGTTGCG GCGCTCGGTT GGTGTGGCCC CTGTGGTGCT TCGTTTCCCC CTCTTTCTCT TTGTTCGGGG 25 GTTCTTGTGG CGGGCTGCTT GTCTCGTTCC GCCCTGTCGG GCGGGAAGCC TCTCTCCTCT CCCCAGATC CGCGACAGGC CGCAGGCAAG AACCAGCGCA ACCAGGGCGC GTCCGCACAG ACTTGGAGGC GGCTGCATGC TGCTACCTGC TCCAGAAGCG TCCGGTGGCC GCCGCGCC CTGTCGGGCG GGBBGCCTCT CTCCTCTCCC CBGBTCCGCG BCBGGCCGCB GGCBBGBBCC BGCGCBBCCB GGGCGCTCC GCBCBGBCTT GGBGGCGGCT GCBTGCTGCT BCCTGCTCGGGCG GGBBGCCTCCG GTGGCCGCCG CGCGTCCGGT GGCCGCCGCG CCTCTCTCCT CTCCCCGTGG CCCTGTCGGG CGGGTCCTGC CGTCCTGTCT. 30 CCTTTTCTTT TGCTGTCTTG TCTTCCCGTC TCTGCTTT GTCTGTCCTC CCCGTCTCCT CCCACTGCTT CTCCCGGGGG GGGTCCGCGC GGCGCCTGGG CCCTTGTGCT GCTTTTTGCT TGTTCCGTTC TGGCTGCTCC GGTCTGTGTT GTGGTTGTTT TGTTTCTTCT TGGGTGTGGG CCTTGCGGTT TTGGCTGTGG GCCCTTTGGG GCCTTGGCTT CTGGCTCGTC TGTCCTCCCC GTCTCCTCCC ACTGCTTCT CCCGGGGGCT TCCCCGGCTT CGGGTGGCCG GTGTCCCGGG CTCCGGCGCG GCGGCGGCTT 35 CGGCTGCGGG TGGGTGGCGC GGGCTGCCGG GTCCGCGCGG CGCCTGGGCC CTTGTGCTGC TTTTTGCTTG TTCCGTTCTG GCTGCTCCGG TCTGTGTTGT GGTTGTTTTG TTTCTTCTTG GGTGTGGGCC TTGCGGTTTT GGCTGTGGGC CCTTTGGGGC CTTGGCTTCT GGCTCCAT CCACATGATT GCTTAGATTT GTGCTGTATC TCTCAGGATT ATCACTGATT ACACATCCAA CCAGTGCCAG CCAAAAGGAT GCCCTGAGGC AAAGGGTTTC CATCTTGAGG CAAATTTGAG GACBTCCBC BTGBTTGCTT BGBTTTGTGC TGTBTCTCTC BGGBTTBTCB CTGBTTBCBC BTCCBBCCBG TGCCBGCCBB BBGGBTGCCC TGBGGCBBBG 40 GGTTTCCBTC TTGBGGCBBB TTTGBGGBGGGCTBBGBT GBTCCBCBTC BCTBCCBCGT TGCCCBCCBC BGBGGTCBCC BCBBTGBCCG TGTBGGCBGC TGCCCBBBGG BCBBTTTGCC BGGCTGGTTG CBCGBBCTGB TTGGGTTCCG BGGTGTTBGT GGBGBTGTTT GGGGBGBGGT CTGBGTCCBC CGGGBGGBCG TTBTCCBTTT CGBBGCTBGG CGGTBBBGCC CTBCTBTCTG TBCBCBBCCC CCCTCTGCBG CBGBGTCCTG TCGTGGCGCC TGGGGCTCBG GGTCCGGGC TAAGATGATC CACATCACTA CCACGTTGCC CACCACAGAG GTCACCACAA TGACCGTGTA GGCAGCTGCC CAAAGGACAA TTTGCCAGGC TGGTTGCACG 45 AACTGATTGG GTTCCGAGGT GTTAGTGGAG ATGTTTGGGG AGAGGTCTGA GTCCACCGGG AGGACGTTAT CCATTTCGAA GCTAGGCGGT AAAGCCCTAC TATCTGTACA CAACCCCCCT CTGCAGCAGA GTCCTGTCGT GGCGCCTGGG GCTCAGGGTC CGTCCTGTCG TGGCGCCTGG GGCTCTTCTT TTGTGGGCTC TTTGGTGGCT GTGGCTGTGG TCTCTGTGGT TGCTGCCCTG GGTCTGGGGG TGTGGCCTTG GGGCCGTCCT CTGGCTCCTC CTCGTGGGCC CCC GGTGBCBTTG BGCBTGTCGG CGCGGTCCCG TTBBGBGTGG GCCCGCCAGC CCAGCCACTC CACTTGGGGG CGGGTGGCCA GCACGAACAG CACCCAGAGG 50 AAGGGGGGCG GCCCAGAAGG GCAGCCCGCA GGCCAGGATC AGGTCTGCTG CGGCCGGAGA TAATGGCATT CACCACGCGG CGGCCCAGCG CACGCCGCGC ATCCGGCCCG GGTTCTGACC TGCAGCCCCC GTCTCCTTGG CATTCCTGGG CCCCAGTCAC TCCTCTCCCT GCCCCCCTTG CTGGGGCAGG GACGGGGTG BCBTTGBGCB TGTCGGCGCG GTCCCGTTBB GBGTGGGCCC GCCAGCCCAG CCACTCCACT TGGGGGCGGG TGGCCAGCAC GAACAGCACC CAGAGGAAGG GGGGCGGCCC AGAAGGGCAG

CCCGCAGGCC AGGATCAGGT CTGCTGCGGC CGGAGATAAT GGCATTCACC ACGCGGCGGC CCAGCGCACG CCGCGCATCC GGCCCGGGTT CTGACCTGCA GCCCCCGTCT CCTTGGCATT CCTGGGCCCC AGTCACTCCT CTCCCTGCCC CCCTTGCTGG GGCAGGGACG GCCGTGTTGT CBGTGGTGCT GCCCGTTTGB GGTBTGGCGC TCCBCCBBTT CCCTTTTCTC CTTGTTTTCC GTITCTCTTG CCGTCTGTGG TT-3' (SEQ. ID NO:2409)

Human Enzyme-related Antisense Polynucleotide 5

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5'-CTT GCT CCT GGG GGC CTC CTG GTC CCT CTG GCT G TT CCC GGC CCT GGB CTG GGG CBG GGG CCG CGT BGG CGC CTG GCT GT GCC CTG GGG GTC TGG GTT CGG CTG T CCC CBG CBG GBC CBG TCC CBT CCB CBG CGT GTG BTG BGT BGC CBT TCT CCT GCB GCC GBG GGG CGC GGG CGB GCB TCG C TTT GGG CTT TTC TCC TTT GGT T TGB GCG CCB GGB CCG CGC BCB GCB GCB GGG CGC GGG CGB GCB TCG CBG CGG CGG GCB GGG GGGCTCCCGC CGCGBGBGGT TBTGGGCTCC CBGGBCCBCC CGCBCCGCGC GGBCGTTTBC BTTCGCCBCG CBGTGCGCGG CCGBCBTGBC GBBGTTGGGC GCBBTCBGGG TGGCGCCGCB GBBGTGGCCT CCGCGCBGCT GCBGGGBCBC CBTGBBGGGC CBCGCGTGGG GCCGCGCTCG CCGGCCCCCC BCBBTCTCCG BGGCCBGCGC GGTGCCCCCC BGCBGCBBGG CCGGCBGGBC BCBGGCGBGG BGBCBCGCGB GTCGGCGGCC GBGGGTCBTG GTGGGGCTGG GGCTCCGGGG TCTCTGCCCC TCCGTGCTGG TGGGGCTGGG GCTCCGGGG TCTCTGCCCC TCCGTGCCGC GTGGGGCCGC GCTCGCCGGC CCCCCCTGC CGGGTGGGCT CCCGCCGCGC GCCGGCCTGC CGGCCCTCG TGGGTCCTGC TGGCCGGGTC CGGGTCCCGG GGGTGGGGCG CGBGTCGGCG GCCGBGGGTC CCCTCCBCBT CTGCTCTGBC CTGCTGGBCT CTGGBTCTGB BGBTBCGCCB TGTBGGGGCG GGBGTGGGGC CTGCTCTCCC GGCCTCCGBT GBTCTCCCCT GCCTCBGCCC CBGTGGGTBG GBGBBGGCC BGCBGBBGCB GGBGTGGCTG CBTCTTTCCT GGTGGGGCCT GCTCTCCCGG CCTGGBGTGG CTCBGCCTGG GCCTGCBGGG CCBCCBGGBG BBTGGCBGCB BGGBTGGCGB GGGTCCTCBT GGCTGGGGTC BCBGBTCCTC TBGCTBGGCB GGGTGBCCBG BGBGGGC GGG TCC TCB TGG CTG GGG GCC TGC BGG GCC GCT CTT GCC TGG BGT GGC TC GCC CBG BGT CTT CCC TGG T CGCTGCBBTC TGCTCCGGGG CTGCBGCBBC CTCBTCBGCTC TTGCCTGGBGTG GCTCBGCCTGG GCCTGCBGGG CCBCCBGGBGB BTGGCBGCBBGG GBTGGCGBGGG TCCTCBTGGC TGGGGTCBCCT GGBGGBGGGG GBGCBGGGGG TCCTCBTGGC TGGGGTCCCT CTCTCCCGTC CT CGG TTT CCT TTG CGG CCT CTG CCC GTG CTC GCCCTG CCT GGG CTG GCC TCT TCG GGT GTG GCT TTG GGG CTC TCT TGG TTG CCC TTT CTT TCGGGTGGGC CGGTGGTGBG CGGCGGCGBCB CGCGGBBGGC CCTGCGCGCC GBGBTCBCCTG CBGGGBGBBG TBGGCTTGCB GCBGGBCTCC CBGGBGGGTG BCBGCBGCCB GTBGBGCTBC CTCGTCCTTC BTGGTBCCGT CGGTGTGGTG GCBCGGGCTG TGTGTGBBGG CGBGCTGGGC CCCGTCTGCT GCTCCTCGTG CCGCCTCGTC CTTCA TGG TA CCGTCGGTGT GGTGGCCTCG GGTGGGCCGG TGGTGGGCG CGCGCGCTCG CGTGGCTCCG GCTCTTCTTT CCCGGCTCCGT CGGCCCGGGG GCCTTGGTCT CCCTCGTCCT TCBTGGTBCC G BCCGGCGGBG CCGCCBGGGT GGBCTGGGBG TGGGTTTCTC CCCGCCGTTC TCBCCCBCCG CGCTGBGCTC BGCGCCTBBG BCTGCTGTTT CTGGBGCTCC TTGGCBBGCC BCBBBCBGCB GBGBGBBBBT CBTGBGCBBB TBBTCCBTTC TGBBBBBBBG GGBTCBBBBB CCTCCCGTTC CCCGTTCGCC TGGCGCGCG TGCGGGTTCC TCGTGGGTTT CTCCCCGCCG TICTCCGGTC TGTTGCCTTT GTGGGCTTCT TGTCTTTTTG GCTGTTCTTT TCCTGCTTGG CGTCTTTTCC TTTCTTTGTG CTCGGTTGTG GGTCCGCTGG TCCTTTGCCC TGTGTGTTTC TGCTGCCCGT TCGCCTGGCG CGCGCTGCGG GTTCCTCGTG GGTTTCTCCC CGCCGTTCTC CGGTCTGTTG CCTTTGTGGG CTTCTTGTCT TTTTGGCTGT TCTTTTCCTG CTTGGCGTCT TITCCTTTCT TTGTGCTCGG TTGTGGGTCC GCTGGTCCTT TGCCCTGTGT GTTTCTGCTG GGBGCTGBTB CTGCBGATTT CBGBGGGBBG BBCCCTGBTB CTCBCCBGCT TCBGCTCTGG BGCBCBBGBG BBBGBGCBGC BGGGGGBGBG GBBGBBGCBG CETCTTCCCB GBGBGGCTGC CTGBGCBBBT GCTGGTTTTC CTTTCCBGTC TTGGGTTTTB TBBCTCCCBG CTTCCTTGTT CCTGGGGGTG TCCTCTTGCT CTGGGCTTTT CTCCCCTTTT CCTTCCTGTC TGTTTTCCTG GGGCTCTCCT CTGTCTCTGT GTCCTTGCCC TGGCCCTCTT CCCTCTCCTG TCTCCTGTCC CTGTGTTCCG CCCGTCTTCC CTCTCCTGAC 50 CTCCTTTTCC TCCGCTGGGT GGGGCCCTGC CTGTTCTCTG CTCCCTGGCT TGGGGTTTCT TCTGTGTGTC TTCTTCCTCT GTTGGCTGGC TTTCTCCTTC TTTTGTCTTC CTGGGTGCCC CTTCTTCCTT TCTTGGGTCC TTGGTGCTTG GGCTGGG

GCGTCTTGGG GTGCBGGGCC CBTCCTGCTG CGCCTGGGCG CTGCTGTGCG TCCGTCTGCT GGGGGGCCGG GGTGGCTGGG CCCTGCTTGC CGCACGACCC CGGGCCGACC CGAGGCTCGG GGGGCTGTGT TCTGGCGCTG GTGGGCTTGG GCCCCTCTGG GGGCTGGGTT TCCTGCTGCG CCTGGGCGCT GGCGTCTTGG GGTGCGGGGC CGGGGGGCCG GGGGGCCGCT GTTCGTGGGC TTTCCTGGGG TCCGCGTGGG GTGCTCCGGT TCCTCGTGCC GCTGCTGCCT TGTCTTTCCG GCCGTGGCGG CGTGGTGGTC 5 CGCCCCCCT GGCCTTCTGC TCGGGGTCTG GCTGGTTGCC GGTGCCCTTG GCGGCGGTCT TCTTCCTGGT GGCTCTGGGC CCGGCCGGTC TCGGGCGTCT CGTGTTCGCT CTTGTGCTGT TCCGGCCGCT CCTTCCTCTT CCGCCGCCGC CGCTCCCCGC CCGCTCGTCG CCCTGGCCCG GCCTCCTCCT GGCCGCTGTC TCGGGCGGCG GCCTTGGCGC TCCGTTTGGG GCTGCCTCTG GCGCTTCCGG CCCTCGGCCT GGGCGCTCTC TTCCGCCTGT GCTGGTGGCC CTCGTGGCC CCTCCTGGCC TCCGGTGTCC TGTGGTCCCC CGGCTGGTGG CCGGGCCGGT TGGGCGGCGC TGGGCGCCGG CGGGTCCTCC GGGCTGCCCT TCTCCGCCGG 10 GGGTCCCGCG CTCCTGCTGT TCCCTGGGCT CTTCTGCCTC TCTCCTGGGT GGGTGCTGGG TGCCGGGGTC TCCGGGGCTTG CCCCGCGCTG CTGGGCCGTTC TGCGGTCTTG GGGTTGTCTG TGGCCCCGCT CGTGTCGCCC TCCGTCGCCC GTCGCCGGCC TCGTCCCCTC CTGGGTGCGC GGCGGGCTGG TCCTGGCGTT TTGCTCCTTC CTGGGCGTCT TGGGGTGCBG GGCCCBTCCT GCTGCGCCTG GGCGCTGCTG TGCGTCCGTC TGCTGGGGGG CCGGGGTGGC TGGGCCCTGC TTGCCGCACG ACCCCGGGCC GACCCGAGGC TCGGGGGGCT GTGTTCTGGC GCTGGTGGGC TTGGGCCCCT CTGGGGGCTG GGTTTCCTGC TGCGCCTGGG 15 CGCTGGCGTC TTGGGGTGCG GGGCCGGGGG GCCGGGGGGC CGCTGTTCGT GGGCCTGGGG GTGCCTGTGG CTGCCGGTTG CCCCGGTTGG TGGCGCCGTC CTGCTGCCGG TCGTTGGCTG GGTCCCCCCG CCCGTTTCCT GGGGTCCGCG TGGGGTGCTC CGGTTCCTCG TGCCGCTGCT GCCTTGTCTT TCCGGCCGTG GCGGCGTGGT GGTCCGCCCC CCCTGGCCTT CTGCTCGGGG TCTGGCTGGT TGCCGGTGCC CTTGGCGGCG GTCTTCTTCC TGGTGGCTCT GGGCCCGGCC GGTCTCGGGC GTCTCGTGTT 20 TCCTGGCCGC TGTCTCGGGC GGCGGCCTTG GCGCTCCGTT TGGGGCTGCC TCTGGCGCTT CCGGCCCTCG GCCTGGGCGC TCTCTTCCGC CTGTGCTGGT GGCCCTCGTG GGCCCCCCTCGT GGCCTCGGT GTCCTGTGGT CCCCCGGCTG GTGGCCGGGC CGGTTGGGCG GGCGTGGGCG CCGGGGGGTC CTCCGGGCTG CCCTTCTCCG CCGGGGGTCC CGCGCTCCTG CTGTTCCCTG GGCTCTTCTG CCTCTCCCT GGGTGGGTGC TGGGTGCCGG GGTCTCCGGG CTTGCCCCGC GCTGCTGGGC GTTCTGCGGT CTTGGGGTTG TCTGTGGCCC CGCTCGTGTC GCCCTCCGTC GCCCGTCGCC GGCCTCGTCC CCTCCTGGGT GCGCGGCGGG 25 CTGGTCCTGG CGTTTTGCTC CTTCCTGG CTGCCCCBGT TTTTGBTCCT CBCBTGCCGT GGGGBGGBCB BTGGCTGCCT CCCCGGGGTT TCTGCTGCTT GCTGCTTCTT TCCCGTCTCC CTTCTTTCCC GTCTCCTTTT TGCCTCTTTG GGTTCCTGTT GTTTCTGGCC TGCTTGGTGG CGGCTTGTGC GTTTCCTCTC TCTTCTCTTG GGTCTCCGCT TCTCGTCCTG CCTTTTCCTG TCTCTGTCGC GCCGTTCCTC CTCCGGCGTC CTCCTGCCCT GTGCTGTTTG CCTCGGGTGG TGCGGGTCCC GGTGCTCCCC CGGCGGCCG GCTGGTTGCC TGGGCCTGTC TGGTGGGGTG TGGGGCCGCT GGGTTGGGGG TGTGGTGGGC TCTTCTGTGG 30 CCTGTGGGGC TGTTGGGGCG TGTGCTGGGT CTTGGGGCTT CCTCCCTTGT GCTGGGTGCG GCCTCCCCGC CCCCCTTCTG GGCCGGTGGC CTGGCTCCTT GTGGGCGCTT CTGGCTCTTG CCCTGTCCTT CTTCGCCTCG TGGCTGCTGG GCTGC-3' (SEQ. ID NO:2410)

Human Factor Related Anti-sense Oligonucleotide

5'-CCT CCT TCC TGG TCT GTC TGC CBG BCB BBT TTG GGB BGT GBB CBG TTT TGG BBC CBT GTT TCC CBG TCT CTG BGC 35 CTT CTG TCC C TGT TTG CTG GTG TCT GCG C 5'- CCC CBB CBG BBG BBG CBG BCB BBT TTG GGB BGT GBB CBG TTT TGG CC CTC TGT TCT TGT TTT GGG GGC GGG CCC GGC CGT TGT CTT G GTT TGG GGG TTT CCG TTG GGG TTC TCC TGG 40 TTT GGT G 5'-GCB CCG TCC BGT GBT GGT GCG GTB CTT GTC GCT GCB GCG CTC GGC CTG GTC CCG GBG BGC GCG CGG GGC TCT GGT TCC CC GCT GCG CCC GTT GTC CTC TGG GGT GGC CTT C GCT CCC GGG TCT GGT TCT TGT GT TGG GGG CCC GGG GCB GGB TGB CTT TTG BGG GGG BCB CBG BTG TCT GGG CBT TGC CBG GTC CTG GGB BCB GBG CCC CGB GCB 45 GGB CCB GGB GTG CGG GCB GCG CGG GCC GGG GGC TGC TGG GBG CCB TBG CGB GGC TGB G CCT CTT TTC TGT TTT TIT GIT GIT TIC TCT TCC TIG CTG BGC BBG BTB TCT BGB TTC TGG GGT GGT CTC GBT TIT BBBB GCT TGB GBB GCT GCB BBC BTT BTC CBB BGT BTB TTT GBG GCT CCB BGG BTC BCG BCC BTC TTC CCB GGC BTT TTB BGT TGC TGT CGT BBG TGB GBG CTG BGB GBB BCT GTG BBG CBB TCB TGB CTT CBB GBG TTC TTT TCB CCC GTT CTT GGC TTC TTC TGT 50 CTG G GTT CTT GGC TTC TGT CCG T TGG CTT CTC GTT GTC CC TGT GGG CTT CTC GTT GTC CC CCC TTC GGG

TIT TIT CIC CIT CIT CIC TCC TIT CIT TIC TIT TCT CIT TCG CIT TCT TIT CGT CIC CIG TIC CIC CIT TT TTG CIG TIT TIT CTC CTT CTT CTC TCC TTT CTT TTC CTC TGT CTT GTT CTG GTC CTT CGT GGG GCT CTG TGT CGC GTG G GTG CGG CCG TGG CC GGC GGB CCB GGB GTT GGB GCB GGB GCB GGB GGB GGC GGC TCB TGT TTG GBT CGG CBG GBG GCB CTC CTC TGG TTG GCT TCC TTC GCC GGC BCB TGC TBG CBG GBB GBB CBG BGG GGG BBG CBG TTG GGB GGT GBG 5 BCC CBT TBB TBG GTG TCG B TCCCTGTTTC CCCCCTTTCG TTCTGCGTTT GCCTTTGGCG TTTTTTGTTT GTTTTCTCTC TCCGTCTTTC TTCTCCCCT GTGGGBBTTT CTGTGGGGBT GGCBTBCBCG TBGGCBGCTC CBBGBGCTBG CBBBCTCBBB TGCBGBBGCB TCCTCBTGGC TCTGBBBCGG TGGGAATTTC TGTGGGGBTG GCATACACGT AGGCAGCTCC AAGAGCTAGC AAACTCAAAT GCAGAAGCATC CTCATGGCTC TGAAACG GGGGGTGGCT TCCTGCCGCG TCTCTGGGCC GTCCCGTCCC 10 CTGCGGGCGC TGTCTCCTGG CTTGTCTTCC GGCTCTTCTG CTGGGGTGGG GCTGGGCGGC CGGCCCGGTG CTGGGGCTCC TCGGGGGGGG GGCTCTTCC GGCCTGTCTC CCTCCGGGGC GGGGTTTCT GGCCGTGGGG GTCTTGCCTG GCCTCCGGGC TECTGCTTGT CTTGCCTTCC TTCTCTGGTC GGTTGTGGCT CGGGGCTCCG TGGGTCCCTG GCGCCCGTTT GTGTTTTGTC 15 TTGGTTTTGG GCTTTTTTC TCTTCCTCCT TTTTCGTGCG TGGGCCTCC GCACGCCTCT TGCCACCTCC TGCGCAGGGC AGCGCCTTGG GGCCAGCGCC GCTCCCGGCG CGGCCAGCAG GGCAGCCAGC AGCGCGAGC CGACGGCCAG CATGCTTCCT CCTCGGCTAC CACTCCATGG TCCCGCAGAG GCGGACAGGC GCBCGCCTC TTGCCBCCTC CTGCGCBGGG CBGCGCCTTG GGGCCBGCGC CGCTCCCGGC GCGGCCBGCB GGGCBGCCBG CBGCGCGCGC GCBCGGCCB GCBTGCTTCC TCCTCGGCTB CCBCTCCBTG GTCCCGCBGB GGCGGBCBGG C GGGGTGGBBB GGTTTGGBGT BTGTCTTTBT GCBCTGBCBT CTBBGTTCTT 20 TBGCBCTCCT TGGCBBBBCT GCBCCTTCBC BCBGBGCTGC BGBBBTCBGG BBGGCTGCCB BGBGBGCCBC GGCCBGCTTG GBBGTCBTGT TTBCBCBCBG TGBGBTGGTT CCTTCCGGGC TTGTGTGCTC TGCTGTCTCT TGGTTCCTTC CGGTGGTTTC TTCCTGGCTC TTGTCCTTTC TCTTGG CCCT TGGC CGGGBGTGGG GGTCCTGGBC GGCBCTGBBG GCBTCCBGGG CTCCCTTCCB GTCCTTCTTG TCCGCTGCCB GCBCCCCTTC BTTCCBGBGG CTGBTGGCCT CCBCCBGGGB CBTGBTTBGG 25 CTTTCCTCCT GGGGCTGCTG CTGGGCTCTT CTTTTTGTTT CTGGCCTGGT GCTCTCTCGT GCCCTTTCCC TTGGGTGTCT TGTTTTTGTG GCCTCCBCCB GGGBCBTG GTCTTTGTTT CTGGGCTCGT GCCCCBTCCC GGCTTCTCTC TGGTTCCGTC CTCTGTGGTG TTTGGCCCTG CTTCCTTTTG CCTGTTGAGG GGGCAGCAGT TGGGCCCCAA AGGCCCTCTC GTTCACCTTC TGGCACGGAGTT GCATCCCCATA GTCAAACTCT GTGGTCGTGT CATAGTCCTC TGTGGTGTTT GGAGTFTCCA TCCCGGCTTC TCTCTGGTTC CAAGGGAGB GGGGGCBGCB GTTGGGCCCC BBBGGCCCTC TCGTTCBCCT TCTGGCBCGG BGTTGCBTCC 30 CCBTBGTCBB BCTCTGTGGT CGTGTCBTBG TCCTCTGTGG TGTTTGGBGT TTCCBTCCCG GCTTCTCTCT GGTTCCBBGG GB GGGCBCGGGG CBGTGGGCGG GCBBTGTBGG CBBBGCBGCB GGGTGTGGTG TCCGBGGBBT BTGGGGBGGC BGBTGCBGGB GCGCBGBGGG CBGTBGCBBT GBGGBTGBCB GCGBGGCGTG CCGCGGBGBC CTTCBTGGTB CCTGTGGBGB GGCTGTCGGB GGGGGTGTGG TGTCCGCTTG GCGGTTCTTT CGGGTGTTTC TTCTCTGGGT TGGCCTGCTG CTCGTCGTGGT CGCTCCGCTC CCGGGTTCGT CTCGCTCTGT CGCCCCTTCC TTCCTTGTCG TGTTCCTCCC TTCCTTGCCT CT GBTGTTTGTT BCCBBBGCBT 35 CBBGBBTBGC TTTGCTBTCT BBGGBTCBCB TTTBGBCBTB GGBBBBCGCT GTBGGTCBGBB BGBTGTGCTT BCCTTCBCBC BGBGCTGCBG BBBTCBGGBBGG CTGCCBBGBGBG CCBCGGCCBGC TTGGBGTCBT GTTTBCBCBC BGTGBGGTGC TCCGGTGGCT TTTTGCTTGT GTGCTCTGCT GTCTCTG TTC CTTCCGGTGG TTTCTTCTCTG GCTCTTGTCC TTTCTCTTGG CCCTTGGCCC CTTGBGCBGG BBGCTCTGGG GCBGGGBGCT GGCBGGGCCC BGGGGGGTGG CTTCCTGCBC TGTCCBGBGT GCBCTGTGCC BCBGCBGCBG CTGCBGGGCC BTCBGCTTCB TGGGGCTCTG GGTGGCBGGT CCBGCCBTGG GTCTGGGTGG 40 GGCTGGGCTG CBGGCTCCGG GCGGTCCBGCCBTGGGTCTG GGGGCTGGG CTGCBGGCTC CGGGCGGGCG GGTGCGGGCT GCGGGCTGCG TGCTGGGGGC TGCCCCGCAG GCCCTGC GCBCCCCTGT GBGCCCTGGG GCCCCCCTGT CTTCTTGGGG BGCGCCTCCT CGGCCBGCTC CBCGTCCCGG BTCBTGCTTT CBGTGCTCBT GGTGTCCTTT CCBGGGGBGB GBGGGGCTGG TCCTCTGCTG TCCTTGCTGG TGCTCBTGGT GTCCTTTCCG CCCTGGGGCC CCCCTGTCTT CTTGGGGCCT CTTCCCTCTG 45 GGGGCCGTCT CTCTCCCTCT CTTGCGTCTC TCTCTTTCTC TCTCTCTTT CCCCTTTCCC GCTCTTTCTG TCTCGGTGTC TGGTTTTCTC TCTCCGCTGG CTGCCTGTCT GGCCTGCGCT CTTGGCCTGT GCTGTTCCTC CTCCGGTTCC TGTCCTCTCT. GTCTGTCGCC CCCTCTGGGG TCTCCCTCTG GGTGGTGGTC TTGTTGCTTG GGCTGGGCTC CGTGTCTCCB GTGCTCBTGG TGTCCGCTGB GGGBGCGTCT GCTGGCGCTG GTCCTCTGCTGTC CTTGCTGGTG CTCBTGGTGT CCTTTCCGCC CTGGGGCCCC 50 CCCTTTCCCG CTCTTTCTGT CTCGGTGTCT GGTTTTCTCT CTCCGCTGGC TGCCTGTCTG GCCTGCGCTC TTGGCCTGTG CTGTTCCTCC TCCGGTTCCT GTCCTCTCTG TCTGTCGCCC CCTCTGGGGT CTCCCTCTGG CGTGGTGGTC TTGTTGCTTG GGCTGGGCTC CGTGTCTCCB GTGCTCBTGG TGTCCGCTGB GGGBGCGTCT GCTGGC CTGCTGBGGC TTGGGTCTCC GGGCGBTTCT CTGCBGBBGB TGCTCBBBGG GCTCCGGCBG TTCCTCCTTG BTCTGGTCGCT GTCGTBCCBG TCGGBCCBGT

BBTTCBGBTC BTCBTTGGCT CCTBTTTCTT CTGCBBBCBG CTGBGTGGBG BCBBGBBBB BGBCTGCCBB GGCCBCGBGG BTTTTCBTGT TGGBTTTTGC GBCGGBCBGT CCCGCGGGGT GCTGAGTTTC TCTGGTTCCT CCGBGCGCBC GTGGTCGCTC TTTTCCGC CGGCCCTTCT CACTGGAGGC ACCGGGCAGT CCTCCATGGG AGGGTTGGGC TTGGCCGGGG CTGCCCGGTG CCTCCTCTIG GCTGGTCCCT CGTTGTCCTT GGGCCCCGC TCCCGCTGCT CGGCCTCCGT GTTCTTTGGC CTCTTGCTCC GCCTGCTGTC TTGTCCCGTC CCCTCCTCGC TTGCGTTTCC CTCTTCCTTG TCTTCCAGGC CTTCCTCCGC TTCCGCTGCT GGGGCCCGCG CCGGGGGGC GCTCGGCTCC GCGGCTTCCT CCCCGGCTGG GGGGTCCTGG TCTCCGGGGC CTGCGGCTCG CGGGCTCGGG GCTGCGTGCG CCGCGCGGG CGTCCGCGGT GGGTGGCGCT GTCCCGCCGT GGTGTGTCTC CGTTCTCGTC CTGCGCCGTC CTGGTCTGCC CGTGGGGTCC TGGGCGTGGT GGGGGGCGTC TGGTGCCCCGT GGGGCTTCGG GCTCGGGGCT GTTCGTCCCC CCTGCCGCTC TGTGGCCTCC GGGGCTCCTC GTTTTCGCTG CTTCGGGTGT CCTTCTCGGC GTGTGGCCCC GGGTCCCGGC CCTGCTGGGC TGGGCGGGGT CGCTGCCCTG GGCTTCTGGC CCGTCTGGTT GTCTGTCGGT GCTTGTCTCG GGTTTCTGGC CTCTGTGCTG GGCGCTTCTC TGCCTCCTGC TCCGCCCTCC TGGTGGCTCG GCTGGGGGTG CCCGTGCGGG GGTGGGTGTG GGGTGTTTTC GGGGTCCTCC CCTTCCC GTT TCA TCT TGG CTT TAT CCTCT CCC CTT GTT CCGTGTTGTC BGTGGTGCTG CCCGTTTGBG GTBTGGCGCT CCBCCBBTTC CCTTTTCTCC TTGTTTTCCG TTTCTCTTGC CGTCTGTGGT T-3' (SEQ. ID NO:2411)

Human Adenosine A, Receptor Anti-sense Oligonucleotide Fragments 20

5'-GAT GGA GGG CGG CAT GGC GGG-3' (FRAG. NO: 1657) (SEQ ID NO:1670)

5'-G CGG GTC GCC GG-3' (FRAG. NO: 1658) (SEQ ID NO:1671) (SEO ID NO:1672)

5'-GGC GGG CBC BGG C-3' (FRAG. NO: 1659)

5'-GGC GGC CBC-3' (FRAG. NO: 1660) 5'-GC GGC CTG G-3' (FRAG. NO: 1661) (SEQ ID NO:1673) (SEQ ID NO:1674)

25 5'-GGB GGG CGG C-3' (FRAG. NO: 1662) (SEQ ID NO:1675) (SEO ID NO:1676) 5'-GBT GGB GGG-3' (FRAG. NO: 1663)

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(SEQ ID NO:1677) 5'-GG CTG GGC-3' (FRAG. NO: 1664)

5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG.1) (SEQ. ID NO: 11) 5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 2) (SEQ. ID NO: 12) 5'-GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 3) (SEQ. ID NO: 13) 30 5'-GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 4) (SEQ. ID NO: 14) 5'-C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 5) (SEQ. ID NO: 15) 5'-CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 6) (SEQ. ID NO: 16)

5'-TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 7) (SEQ. ID NO: 17) 35 5'-G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 8) (SEQ. ID NO: 18) 5'-GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 9) (SEQ. ID NO: 19) 5'-AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 10) (SEQ. ID NO: 20) 5'-A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 11) (SEQ. ID NO: 21)

5'-AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 12) (SEQ. ID NO: 22) 40 5'-GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 13) (SEQ. ID NO: 23) 5'-C TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 14) (SEQ. ID NO: 24) 5'-TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 15) (SEQ. ID NO: 25)

5'-GA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 16) (SEQ. ID NO: 26) 5'-A GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 17) (SEQ. ID NO: 27) 45 5'-GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 18) (SEQ. ID NO: 28) 5'-AT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 19) (SEQ. ID NO: 29) 5'-T GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 20) (SEQ. ID NO: 30)

5'-GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 21) (SEQ. ID NO: 31) 5'-GA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 22) (SEQ. ID NO: 32) 50 5'-A GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 23) (SEQ. ID NO: 33) 5'-GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 24) (SEQ. ID NO: 34) 5'-GG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 25) (SEQ. ID NO: 35) 5'-G CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 26) (SEQ. ID NO: 36)

5'-CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 27) (SEQ. ID NO: 37) 55 5'-GG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 28) (SEQ. ID NO: 38) 5'-G CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 29) (SEQ. ID NO: 39) 5'-CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 30) (SEQ. ID NO: 40)

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5'-AT GGC GGG CAC AGG CTG GGC-3' (FRAG 31) (SEQ. ID NO: 41)
       5'-T GGC GGG CAC AGG CTG GGC-3' (FRAG 32) (SEQ. ID NO: 42)
       5'-GGC GGG CAC AGG CTG GGC-3' (FRAG 33) (SEQ. ID NO: 43)
       5'-GC GGG CAC AGG CTG GGC-3' (FRAG 34) (SEQ. ID NO: 44)
       5'-C GGG CAC AGG CTG GGC-3' (FRAG 35) (SEQ. ID NO: 45)
5
       5'-GGG CAC AGG CTG GGC-3' (FRAG 36) (SEQ. ID NO: 46)
       5'-GG CAC AGG CTG GGC-3' (FRAG 37) (SEQ. ID NO: 47)
       5'-G CAC AGG CTG GGC-3' (FRAG 38) (SEQ. ID NO: 48)
       5'-CAC AGG CTG GGC-3' (FRAG 39) (SEQ. ID NO: 49)
       5'-AC AGG CTG GGC-3' (FRAG 40) (SEQ. ID NO: 50)
10
       5'-C AGG CTG GGC-3' (FRAG 41) (SEQ. ID NO: 51)
       5'-AGG CTG GGC-3' (FRAG 42) (SEQ. ID NO: 52)
       5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 43) (SEQ. ID NO: 53)
       5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 44) (SEQ. ID NO: 54)
       5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 45) (SEQ. ID NO: 55)
15
       5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 46) (SEQ. ID NO: 56)
       5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 47) (SEQ. ID NO: 57)
       5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 48) (SEQ. ID NO: 58)
       5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 49) (SEQ. ID NO: 59)
       5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 50) (SEQ. ID NO: 60)
20
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 51) (SEQ. ID NO: 61)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 52) (SEQ. ID NO: 62)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 53) (SEQ. ID NO: 63)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 54) (SEQ. ID NO: 64)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 55) (SEQ. ID NO: 65)
25
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 56) (SEQ. ID NO: 66)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3'(FRAG 57) (SEQ. ID NO: 67)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 58) (SEQ. ID NO: 68)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 59) (SEQ. ID NO: 69)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 60) (SEQ. ID NO: 70)
30
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 61) (SEQ. ID NO: 71)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 62) (SEQ. ID NO: 72)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 63) (SEQ. ID NO: 73)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 64) (SEQ. ID NO: 74)
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG CG -3' (FRAG 65) (SEQ. ID NO: 75)
 35
        5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG C -3' (FRAG 66) (SEQ. ID NO: 76)
         5'-GGC GGC CTG GAA AGC TGA GAT GGA GGG -3' (FRAG 67) (SEQ. ID NO: 77)
         5'-GGC GGC CTG GAA AGC TGA GAT GGA GG -3' (FRAG 68) (SEQ. ID NO: 78)
         5'-GGC GGC CTG GAA AGC TGA GAT GGA G -3' (FRAG 69) (SEQ. ID NO: 79)
         5'-GGC GGC CTG GAA AGC TGA GAT GGA -3' (FRAG 70) (SEQ. ID NO: 80)
 40
         5'-GGC GGC CTG GAA AGC TGA GAT GG -3' (FRAG 71) (SEQ. ID NO: 81)
         5'-GGC GGC CTG GAA AGC TGA GAT G -3' (FRAG 72) (SEQ. ID NO: 82)
         5'-GGC GGC CTG GAA AGC TGA GAT -3' (FRAG 73) (SEQ. ID NO: 83)
         5'-GGC GGC CTG GAA AGC TGA GA-3' (FRAG 74) (SEQ. ID NO: 84)
         5'-GGC GGC CTG GAA AGC TGA G-3' (FRAG 75) (SEQ. ID NO: 85)
 45
         5'-GGC GGC CTG GAA AGC TGA-3' (FRAG 76) (SEQ. ID NO: 86)
         5'-GGC GGC CTG GAA AGC TG-3' (FRAG 77) (SEQ. ID NO: 87)
         5'-GGC GGC CTG GAA AGC T-3' (FRAG 78) (SEQ. ID NO: 88)
         5'-GGC GGC CTG GAA AGC-3' (FRAG 79) (SEQ. ID NO: 89)
         5'-GGC GGC CTG GAA AG-3' (FRAG 80) (SEQ. ID NO: 90)
 50
         5'-GGC GGC CTG GAA A-3' (FRAG 81) (SEQ. ID NO: 91)
         5'-GGC GGC CTG GAA-3' (FRAG 82) (SEQ. ID NO: 92)
         5'-GGC GGC CTG GA-3' (FRAG 83) (SEQ. ID NO: 93)
         5'-GGC GGC CTG G-3' (FRAG 84) (SEQ. ID NO: 94)
         5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 85) (SEQ. ID NO: 95)
  55
         5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 86) (SEQ. ID NO: 96)
         5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 87) (SEQ. ID NO: 97)
         5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 88) (SEQ. ID NO: 98)
         5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 89) (SEQ. ID NO: 99)
          5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 90) (SEQ. ID NO: 100)
  60
          5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 91) (SEQ. ID NO: 101)
          5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 92) (SEQ. ID NO: 102)
          5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 93) (SEQ. ID NO: 103)
          5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 94) (SEQ. ID NO: 104)
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5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 95) (SEQ. ID NO: 105)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 96) (SEQ. ID NO: 106)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 97) (SEQ. ID NO: 107)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 98) (SEQ. ID NO: 108)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 99) (SEQ. ID NO: 109)
5
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 100) (SEQ. ID NO: 110)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 101) (SEQ. ID NO: 111)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 102) (SEQ. ID NO: 112)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 103) (SEQ. ID NO: 113)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 104) (SEQ. ID NO: 114)
10
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 105) (SEQ. ID NO: 115)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 106) (SEQ. ID NO: 116)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG CG -3' (FRAG 107) (SEQ. ID NO: 117)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG C -3' (FRAG 108) (SEQ. ID NO: 118)
       5'-GC GGC CTG GAA AGC TGA GAT GGA GGG -3' (FRAG 109) (SEQ. ID NO: 119)
15
       5'-GC GGC CTG GAA AGC TGA GAT GGA GG -3' (FRAG 110) (SEQ. ID NO: 120)
       5'-GC GGC CTG GAA AGC TGA GAT GGA G -3' (FRAG 111) (SEQ. ID NO: 121)
        5'-GC GGC CTG GAA AGC TGA GAT GGA -3' (FRAG 112) (SEQ. ID NO: 122)
        5'-GC GGC CTG GAA AGC TGA GAT GG -3' (FRAG 113) (SEQ. ID NO: 123)
        5'-GC GGC CTG GAA AGC TGA GAT G -3' (FRAG 114) (SEQ. ID NO: 124)
20
        5'-GC GGC CTG GAA AGC TGA GAT -3' (FRAG 115) (SEQ. ID NO: 125)
        5'-GC GGC CTG GAA AGC TGA GA-3' (FRAG 116) (SEQ. ID NO: 126)
        5'-GC GGC CTG GAA AGC TGA G-3' (FRAG 117) (SEQ. ID NO: 127)
        5'-GC GGC CTG GAA AGC TGA-3' (FRAG 118) (SEQ. ID NO: 128)
        5'-GC GGC CTG GAA AGC TG-3' (FRAG 119) (SEQ. ID NO: 129)
25
        5'-GC GGC CTG GAA AGC T-3' (FRAG 120) (SEQ. ID NO: 130)
        5'-GC GGC CTG GAA AGC-3' (FRAG 121) (SEQ. ID NO: 131)
        5'-GC GGC CTG GAA AG-3' (FRAG 122) (SEQ: ID NO: 132)
        5'-GC GGC CTG GAA A-3' (FRAG 123) (SEQ. ID NO: 133)
        5'-GC GGC CTG GAA-3' (FRAG 124) (SEQ. ID NO: 134)
30
        5'-GC GGC CTG GA-3' (FRAG 125) (SEQ. ID NO: 135)
        5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 126) (SEQ. ID NO: 136)
        5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 127) (SEQ. ID NO: 137)
        5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 128) (SEQ. ID NO: 138)
        5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 129) (SEQ. ID NO: 139)
 35
        5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 130) (SEQ. ID NO: 140)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 131) (SEQ. ID NO: 141)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 132) (SEQ. ID NO: 142)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 133) (SEQ. ID NO: 143)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 134) (SEQ. ID NO: 144)
 40
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 135) (SEQ. ID NO: 145)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 136) (SEQ. ID NO: 146)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 137) (SEQ. ID NO: 147)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 138) (SEQ. ID NO: 148)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 139) (SEQ. ID NO: 149)
 45
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 140) (SEQ. ID NO: 150)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 141) (SEQ. ID NO: 151)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 142) (SEQ. ID NO: 152)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRA 143) (SEQ. ID NO: 153)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 144) (SEQ. ID NO: 154)
  50
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 145) (SEQ. ID NO: 155)
         5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 146) (SEQ. ID NO: 156)
          5'-C GGC CTG GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 147) (SEQ. ID NO: 157)
          5'-C GGC CTG GAA AGC TGA GAT GGA GGG CG -3' (FRAG 148) (SEQ. ID NO: 158)
          5'-C GGC CTG GAA AGC TGA GAT GGA GGG C -3' (FRAG 148) (SEQ. ID NO: 159)
  55
          5'-C GGC CTG GAA AGC TGA GAT GGA GGG -3' (FRAG 150) (SEQ. ID NO: 160)
          5'-C GGC CTG GAA AGC TGA GAT GGA GG -3' (FRAG 151) (SEQ. ID NO: 161)
          5'-C GGC CTG GAA AGC TGA GAT GGA G -3' (FRAG 152) (SEQ. ID NO: 162)
          5'-C GGC CTG GAA AGC TGA GAT GGA -3' (FRAG 153) (SEQ. ID NO: 163)
          5'-C GGC CTG GAA AGC TGA GAT GG -3' (FRAG 154) (SEQ. ID NO: 164)
  60
          5'-C GGC CTG GAA AGC TGA GAT G -3' (FRAG 155) (SEQ. ID NO: 165)
          5'-C GGC CTG GAA AGC TGA GAT -3' (FRAG 156) (SEQ. ID NO: 166)
          5'-C GGC CTG GAA AGC TGA GA-3' (FRAG 157) (SEQ. ID NO: 167)
          5'-C GGC CTG GAA AGC TGA G-3' (FRAG 158) (SEQ. ID NO: 168)
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5'-C GGC CTG GAA AGC TGA-3' (FRAG 159) (SEQ. ID NO: 169)
       5'-C GGC CTG GAA AGC TG-3' (FRAG 160) (SEQ. ID NO: 170)
       5'-C GGC CTG GAA AGC T-3' (FRAG 161) (SEQ. ID NO: 171)
       5'-C GGC CTG GAA AGC-3' (FRAG 162) (SEQ. ID NO: 172)
       5'-C GGC CTG GAA AG-3' (FRAG 163) (SEQ. ID NO: 173)
 5
       5'-C GGC CTG GAA A-3' (FRAG 164) (SEQ. ID NO: 174)
       5'-C GGC CTG GAA-3' (FRAG 165) (SEQ. ID NO: 175)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 166) (SEQ. ID NO: 176)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 167) (SEQ. ID NO: 177)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 168) (SEQ. ID NO: 178)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 169) (SEQ. ID NO: 179)
10
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 170) (SEQ. ID NO: 180)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 171) (SEQ. ID NO: 181)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 172) (SEQ. ID NO: 182)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 173) (SEQ. ID NO: 183)
15
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 174) (SEQ. ID NO: 184)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 175) (SEQ. ID NO: 185)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 176) (SEQ. ID NO: 186)
       5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 177) (SEQ. ID NO: 187)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 178) (SEQ. ID NO: 188)
20
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 179) (SEQ. ID NO: 189)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 180) (SEQ. ID NO: 190)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 181) (SEQ. ID NO: 191)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 182) (SEQ. ID NO: 192)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 183) (SEQ. ID NO: 193)
25
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 184) (SEQ. ID NO: 194)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 185) (SEQ. ID NO: 195)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 186) (SEQ. ID NO: 196)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 187) (SEQ. ID NO: 197)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG CG -3' (FRAG 188) (SEQ. ID NO: 198)
 30
        5'- GGC CTG GAA AGC TGA GAT GGA GGG C -3' (FRAG 189) (SEQ. ID NO: 199)
        5'- GGC CTG GAA AGC TGA GAT GGA GGG -3' (FRAG 190) (SEQ. ID NO: 200)
        5'- GGC CTG GAA AGC TGA GAT GGA GG -3' (FRAG 191) (SEQ. ID NO: 201)
        5'- GGC CTG GAA AGC TGA GAT GGA G -3' (FRAG 192) (SEQ. ID NO: 202)
         5'- GGC CTG GAA AGC TGA GAT GGA -3' (FRAG 193) (SEQ. ID NO: 203)
 35
         5'- GGC CTG GAA AGC TGA GAT GG -3' (FRAG 194) (SEQ. ID NO: 204)
         5'- GGC CTG GAA AGC TGA GAT G -3' (FRAG 195) (SEQ. ID NO: 205)
         5'- GGC CTG GAA AGC TGA GAT -3' (FRAG 196) (SEQ. ID NO: 206)
         5'- GGC CTG GAA AGC TGA GA-3' (FRAG 197) (SEQ. ID NO: 207)
         5'- GGC CTG GAA AGC TGA G-3' (FRAG 198) (SEQ. ID NO: 208)
 40
         5'- GGC CTG GAA AGC TGA-3' (FRAG 199) (SEQ. ID NO: 209)
         5'- GGC CTG GAA AGC TG-3' (FRAG 200 (SEQ. ID NO: 210)
         5'- GGC CTG GAA AGC T-3' (FRAG 201) (SEQ. ID NO: 211)
         5'- GGC CTG GAA AGC-3' (FRAG 202) (SEQ. ID NO: 212)
         5'- GGC CTG GAA AG-3' (FRAG 203) (SEQ. ID NO: 213)
 45 .
          5'- GGC CTG GAA A-3' (FRAG 204) (SEQ. ID NO: 214)
         5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 205) (SEQ. ID NO: 215)
         5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 206) (SEQ. ID NO: 216)
         5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 207) (SEQ. ID NO: 217)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 208) (SEQ. ID NO: 218)
  50
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 209) (SEQ. ID NO: 219)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 210) (SEQ. ID NO: 220)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 211) (SEQ. ID NO: 221)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 212) (SEQ. ID NO: 222)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 213) (SEQ. ID NO: 223)
  55
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 214) (SEQ. ID NO: 224)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 215) (SEQ. ID NO: 225)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 216) (SEQ. ID NO: 226)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 217) (SEQ. ID NO: 227)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 218) (SEQ. ID NO: 228)
  60
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 219) (SEQ. ID NO: 229)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 220) (SEQ. ID NO: 230)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 221) (SEQ. ID NO: 231)
          5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 222) (SEQ. ID NO: 232)
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5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 223) (SEQ. ID NO: 233)
       5'- GC CTG GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 224) (SEQ. ID NO: 234)
       5'- GC CTG GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 225) (SEQ. ID NO: 235)
       5'- GC CTG GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 226) (SEQ. ID NO: 236)
       5'- GC CTG GAA AGC TGA GAT GGA GGG CG -3' (FRAG 227) (SEQ. ID NO: 237)
5
       5'- GC CTG GAA AGC TGA GAT GGA GGG C -3' (FRAG 228) (SEQ. ID NO: 238)
       5'- GC CTG GAA AGC TGA GAT GGA GGG -3' (FRAG 229) (SEQ. ID NO: 239)
       5'- GC CTG GAA AGC TGA GAT GGA GG -3' (FRAG 230) (SEQ. ID NO: 240)
       5'- GC CTG GAA AGC TGA GAT GGA G -3' (FRAG 231) (SEQ. ID NO: 241)
       5'- GC CTG GAA AGC TGA GAT GGA -3' (FRAG 232) (SEQ. ID NO: 242)
10
       5'- GC CTG GAA AGC TGA GAT GG -3' (FRAG 233) (SEQ. ID NO: 243)
       5'- GC CTG GAA AGC TGA GAT G -3' (FRAG 234) (SEQ. ID NO: 244)
       5'- GC CTG GAA AGC TGA GAT -3' (FRAG 235) (SEQ. ID NO: 245)
        5'- GC CTG GAA AGC TGA GA-3' (FRAG 236) (SEQ. ID NO: 246)
        5'- GC CTG GAA AGC TGA G-3' (FRAG 237) (SEQ. ID NO: 247)
15
        5'- GC CTG GAA AGC TGA-3' (FRAG 238) (SEQ. ID NO: 248)
        5'- GC CTG GAA AGC TG-3' (FRAG 239) (SEQ. ID NO: 249)
        5'- GC CTG GAA AGC T-3' (FRAG 240) (SEQ. ID NO: 250)
        5'- GC CTG GAA AGC-3' (FRAG 241) (SEQ. ID NO: 251)
        5'- GC CTG GAA AG-3' (FRAG 242) (SEQ. ID NO: 252)
        5'- C CTG GAA AGC TGA GAT GG A GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 243) (SEQ. ID NO: 253)
20
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 244) (SEQ. ID NO: 254)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 245) (SEQ. ID NO: 255)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 246) (SEQ. ID NO: 256)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 247) (SEQ. ID NO: 257)
25
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 248) (SEQ. ID NO: 258)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 249) (SEQ. ID NO: 259)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 250) (SEQ. ID NO: 260)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 251) (SEQ. ID NO: 261)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 252) (SEQ. ID NO: 262)
 30
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 253) (SEQ. ID NO: 263)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 254) (SEQ. ID NO: 264)
        5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 255) (SEQ. ID NO: 265)
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 256) (SEQ. ID NO: 266)
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 257) (SEQ. ID NO: 267)
 35
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 258) (SEQ. ID NO: 268)
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 259) (SEQ. ID NO: 269)
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 260) (SEQ. ID NO: 270)
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 261) (SEQ. ID NO: 271)
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 262) (SEQ. ID NO: 272)
 40
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 263) (SEQ. ID NO: 273)
         5'- C CTG GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 264) (SEQ. ID NO: 274)
         5'- C CTG GAA AGC TGA GAT GGA GGG CG -3' (FRAG 265) (SEQ. ID NO: 275)
         5'- C CTG GAA AGC TGA GAT GGA GGG C -3' (FRAG 266) (SEQ. ID NO: 276)
         5'- C CTG GAA AGC TGA GAT GGA GGG -3' (FRAG 267) (SEQ. ID NO: 277)
 45
         5'- C CTG GAA AGC TGA GAT GGA GG -3' (FRAG 268) (SEQ. ID NO: 278)
         5'- C CTG GAA AGC TGA GAT GGA G -3' (FRAG 269) (SEQ. ID NO: 279)
         5'- C CTG GAA AGC TGA GAT GGA -3' (FRAG 270) (SEQ. ID NO: 280)
         5'- C CTG GAA AGC TGA GAT GG -3' (FRAG 271) (SEQ. ID NO: 281)
         5'- C CTG GAA AGC TGA GAT G -3' (FRAG 272) (SEQ. ID NO: 282)
  50
          5'- C CTG GAA AGC TGA GAT -3' (FRAG 273) (SEQ. ID NO: 283)
          5'- C CTG GAA AGC TGA GA-3' (FRAG 274) (SEQ. ID NO: 284)
          5'- C CTG GAA AGC TGA G-3' (FRAG 275) (SEQ. ID NO: 285)
          5'- C CTG GAA AGC TGA-3' (FRAG 276) (SEQ. ID NO: 286)
          5'- C CTG GAA AGC TG-3' (FRAG 277) (SEQ. ID NO: 287)
  55
          5'- C CTG GAA AGC T-3' (FRAG 278) (SEQ. ID NO: 288)
          5'- C CTG GAA AGC-3' (FRAG 279) (SEQ. ID NO: 289)
          5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 280) (SEQ. ID NO: 290)
          5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 281) (SEQ. ID NO: 291)
          5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 282) (SEQ. ID NO: 292)
  60
          5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 283) (SEQ. ID NO: 293)
          5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 284) (SEQ. ID NO: 294)
          5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 285) (SEQ. ID NO: 295)
          5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 286) (SEQ. ID NO: 296)
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5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 287) (SEQ. ID NO: 297)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 288) (SEQ. ID NO: 298)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 289) (SEQ. ID NO: 299)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 290) (SEQ. ID NO: 300)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 291) (SEQ. ID NO: 301)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 292) (SEQ. ID NO: 302)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 293) (SEQ. ID NO: 303)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 294) (SEQ. ID NO: 304)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 295) (SEQ. ID NO: 305)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 296) (SEQ. ID NO: 306)
10
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 297) (SEQ. ID NO: 307)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 298) (SEQ. ID NO: 308)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 299) (SEQ. ID NO: 309)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 300) (SEQ. ID NO: 310)
       5'- CTG GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 301) (SEQ. ID NO: 311)
15
        5'- CTG GAA AGC TGA GAT GGA GGG CG -3' (FRAG 302) (SEQ. ID NO: 312)
        5'- CTG GAA AGC TGA GAT GGA GGG C -3' (FRAG 303) (SEQ. ID NO: 313)
        5'- CTG GAA AGC TGA GAT GGA GGG -3' (FRAG 304) (SEQ. ID NO: 314)
        5'- CTG GAA AGC TGA GAT GGA GG -3' (FRAG 305) (SEQ. ID NO: 315)
        5'- CTG GAA AGC TGA GAT GGA G -3' (FRAG 306) (SEQ. ID NO: 316)
20
        5'- CTG GAA AGC TGA GAT GGA -3' (FRAG 307) (SEQ. ID NO: 317)
        5'- CTG GAA AGC TGA GAT GG -3' (FRAG 308) (SEQ. ID NO: 318)
        5'- CTG GAA AGC TGA GAT G -3' (FRAG 309) (SEQ. ID NO: 319)
        5'- CTG GAA AGC TGA GAT -3' (FRAG 310) (SEQ. ID NO: 320)
        5'- CTG GAA AGC TGA GA-3' (FRAG 311) (SEQ. ID NO: 321)
25
        5'- CTG GAA AGC TGA G-3' (FRAG 312) (SEQ. ID NO: 322)
        5'- CTG GAA AGC TGA-3' (FRAG 313) (SEQ. ID NO: 323)
        5'- CTG GAA AGC TG-3' (FRAG 314) (SEQ. ID NO: 324)
        5'- CTG GAA AGC T-3' (FRAG 315) (SEQ. ID NO: 325)
        5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 316) (SEQ. ID NO: 326)
30
        5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 317) (SEQ. ID NO: 327)
        5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 318) (SEQ. ID NO: 328)
        5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 319) (SEQ. ID NO: 329)
        5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 320) (SEQ. ID NO: 330)
        5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 321) (SEQ. ID NO: 331)
 35
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 322) (SEQ. ID NO: 332)
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 323) (SEQ. ID NO: 333)
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 324) (SEQ. ID NO: 334)
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 325) (SEQ. ID NO: 335)
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 326) (SEQ. ID NO: 336)
 40
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 327) (SEQ. ID NO: 337)
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 328) (SEQ. ID NO: 338)
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 329) (SEQ. ID NO: 339)
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 330) (SEQ. ID NO: 340)
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 331) (SEQ. ID NO: 341)
 45
         5'- TG GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 332) (SEQ. ID NO: 342)
          5'- TG GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 333) (SEQ. ID NO: 343)
          5'- TG GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 334) (SEQ. ID NO: 344)
          5'- TG GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 335) (SEQ. ID NO: 345)
          5'- TG GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 336) (SEQ. ID NO: 346)
  50
          5'- TG GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 337) (SEQ. ID NO: 347)
          5'- TG GAA AGC TGA GAT GGA GGG CG -3' (FRAG 338) (SEQ. ID NO: 348)
          5'- TG GAA AGC TGA GAT GGA GGG C -3' (FRAG 339) (SEQ. ID NO: 349)
          5'- TG GAA AGC TGA GAT GGA GGG -3' (FRAG 340) (SEQ. ID NO: 350)
          5'- TG GAA AGC TGA GAT GGA GG -3' (FRAG 341) (SEQ. ID NO: 351)
  55
          5'- TG GAA AGC TGA GAT GGA G -3' (FRAG 342) (SEQ. ID NO: 352)
          5'- TG GAA AGC TGA GAT GGA -3' (FRAG 343) (SEQ. ID NO: 353)
          5'- TG GAA AGC TGA GAT GG -3' (FRAG 344) (SEQ. ID NO: 354)
          5'- TG GAA AGC TGA GAT G -3' (FRAG 345) (SEQ. ID NO: 355)
          5'- TG GAA AGC TGA GAT -3' (FRAG 346) (SEQ. ID NO: 356)
  60
          5'- TG GAA AGC TGA GA-3' (FRAG 347) (SEQ. ID NO: 357)
          5'- TG GAA AGC TGA G-3' (FRAG 348) (SEQ. ID NO: 358)
          5'- TG GAA AGC TGA-3' (FRAG 349) (SEQ. ID NO: 359)
          5'- TG GAA AGC TG-3' (FRAG 350) (SEQ. ID NO: 360)
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5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 351) (SEQ. ID NO: 361)
      5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 352) (SEQ. ID NO: 362)
      5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 353) (SEQ. ID NO: 363)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 354) (SEQ. ID NO: 364)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 355) (SEQ. ID NO: 365)
5
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 356) (SEQ. ID NO: 366)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 357) (SEQ. ID NO: 367)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 358) (SEQ. ID NO: 368)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 359) (SEQ. ID NO: 369)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 360) (SEQ. ID NO: 370)
10
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 361) (SEQ. ID NO: 371)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 362) (SEQ. ID NO: 372)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 363) (SEQ. ID NO: 373)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 364) (SEQ. ID NO: 374)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 365) (SEQ. ID NO: 375)
15
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 366) (SEQ. ID NO: 376)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 367) (SEQ. ID NO: 377)
        5'- G GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 368) (SEQ. ID NO: 378)
       5'- G GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 369) (SEQ. ID NO: 379)
        5'- G GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 370) (SEQ. ID NO: 380)
20
        5'- G GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 371) (SEQ. ID NO: 381)
        5'- G GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 372) (SEQ. ID NO: 382)
        5'- G GAA AGC TGA GAT GGA GGG CG -3' (FRAG 373) (SEQ. ID NO: 383)
        5'- G GAA AGC TGA GAT GGA GGG C -3' (FRAG 374) (SEQ. ID NO: 384)
        5'- G GAA AGC TGA GAT GGA GGG -3' (FRAG 375) (SEQ. ID NO: 385)
25
        5'- G GAA AGC TGA GAT GGA GG -3' (FRAG 376) (SEQ. ID NO: 386)
        5'- G GAA AGC TGA GAT GGA G -3' (FRAG 377) (SEQ. ID NO: 387)
        5'- G GAA AGC TGA GAT GGA -3' (FRAG 378) (SEQ. ID NO: 388)
        5'- G GAA AGC TGA GAT GG -3' (FRAG 379) (SEQ. ID NO: 389)
        5'- G GAA AGC TGA GAT G -3' (FRAG 380) (SEQ. ID NO: 390)
30
        5'- G GAA AGC TGA GAT -3' (FRAG 381) (SEQ. ID NO: 391)
        5'- G GAA AGC TGA GA-3' (FRAG 382) (SEQ. ID NO: 392)
        5'- G GAA AGC TGA G-3' (FRAG 383) (SEQ. ID NO: 393)
        5'- G GAA AGC TGA-3' (FRAG 384) (SEQ. ID NO: 394)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3* (FRAG 385) (SEQ. ID NO: 395)
 35
        5'- GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 386) (SEQ. ID NO: 396)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 387) (SEQ. ID NO: 397)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 388) (SEQ. ID NO: 398)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 389) (SEQ. ID NO: 399)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 390) (SEQ. ID NO: 400)
 40
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 391) (SEQ. ID NO: 401)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 392) (SEQ. ID NO: 402)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 393) (SEQ. ID NO: 403)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 394) (SEQ. ID NO: 404)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 395) (SEQ. ID NO: 405)
 45
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 396) (SEQ. ID NO: 406)
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 397) (SEQ. ID NO: 407)
         5'-
            GAA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 398) (SEQ. ID NO: 408)
         5'-
            GAA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 399) (SEQ. ID NO: 409)
             GAA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 400) (SEQ. ID NO: 410)
  50
         5'- GAA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 401) (SEQ. ID NO: 411)
             GAA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 402) (SEQ. ID NO: 412)
         5'-
             GAA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 403) (SEQ. ID NO: 413)
         5'-
             GAA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 404) (SEQ. ID NO: 414)
             GAA AGC TGA GAT GGA GGG CGG C-3' (FRAG 405) (SEQ. ID NO: 415)
  55
         5'- GAA AGC TGA GAT GGA GGG CGG -3' (FRAG 406) (SEQ. ID NO: 416)
             GAA AGC TGA GAT GGA GGG CG -3' (FRAG 407) (SEQ. ID NO: 417)
         5'- GAA AGC TGA GAT GGA GGG C -3' (FRAG 408) (SEQ. ID NO: 418)
          5'- GAA AGC TGA GAT GGA GGG -3' (FRAG 409) (SEQ. ID NO: 419)
          5'- GAA AGC TGA GAT GGA GG -3' (FRAG 410) (SEQ. ID NO: 420)
  60
          5'- GAA AGC TGA GAT GGA G -3' (FRAG 411) (SEQ. ID NO: 421)
          5'- GAA AGC TGA GAT GGA -3' (FRAG 412) (SEQ. ID NO: 422)
          5'- GAA AGC TGA GAT GG -3' (FRAG 413) (SEQ. ID NO: 423)
          5'- GAA AGC TGA GAT G -3' (FRAG 414) (SEQ. ID NO: 424)
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5'- GAA AGC TGA GAT -3' (FRAG 415) (SEQ. ID NO: 425)
      5'- GAA AGC TGA GA-3' (FRAG 416) (SEQ. ID NO: 426)
      5'- GAA AGC TGA G-3' (FRAG 417) (SEQ. ID NO: 427)
      5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 418) (SEQ. ID NO: 428)
      5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 419) (SEQ. ID NO: 429)
      5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 420) (SEQ. ID NO: 430)
5
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 421) (SEQ. ID NO: 431)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 422) (SEQ. ID NO: 432)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 423) (SEQ. ID NO: 433)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3'(FRAG 424) (SEQ. ID NO: 434)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 425) (SEQ. ID NO: 435)
10
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 426) (SEQ. ID NO: 436)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 427) (SEQ. ID NO: 437)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 428) (SEQ. ID NO: 438)
           AA AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 429) (SEQ. ID NO: 439)
15
           AA AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 430) (SEQ. ID NO: 440)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 431) (SEQ. ID NO: 441)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 432) (SEQ. ID NO: 442)
       5'- AA AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 433) (SEQ. ID NO: 443)
          AA AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 434) (SEQ. ID NO: 444)
20
       5'- AA AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 435) (SEQ. ID NO: 445)
       5'- AA AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 436) (SEQ. ID NO: 446)
        5'- AA AGC TGA GAT GGA GGG CGG CA-3' (FRAG 437) (SEQ. ID NO: 447)
           AA AGC TGA GAT GGA GGG CGG C-3' (FRAG 438) (SEQ. ID NO: 448)
        5'-
        5'- AA AGC TGA GAT GGA GGG CGG -3' (FRAG 439) (SEQ. ID NO: 449)
25
        5'- AA AGC TGA GAT GGA GGG CG -3' (FRAG 440) (SEQ. ID NO: 450)
        5'- AA AGC TGA GAT GGA GGG C -3' (FRAG 441) (SEQ. ID NO: 451)
        5'- AA AGC TGA GAT GGA GGG -3' (FRAG 442) (SEQ. ID NO: 452)
        5'- AA AGC TGA GAT GGA GG -3' (FRAG 443) (SEQ. ID NO: 453)
        5'- AA AGC TGA GAT GGA G -3' (FRAG 444) (SEQ. ID NO: 454)
 30
            AA AGC TGA GAT GGA -3' (FRAG 445) (SEQ. ID NO: 455)
        5'- AA AGC TGA GAT GG -3' (FRAG 446) (SEQ. ID NO: 456)
            AA AGC TGA GAT G -3' (FRAG 447) (SEQ. ID NO: 457)
            AA AGC TGA GAT -3' (FRAG 448) (SEQ. ID NO: 458)
        5'-
            AA AGC TGA GA-3' (FRAG 449) (SEQ. ID NO: 459)
            A AGC TGA GAT GGA GGG CG G CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 450) (SEQ. ID NO: 460)
 35
            A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 451) (SEQ. ID NO: 461)
            A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 452) (SEQ. ID NO: 462)
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 453) (SEQ. ID NO: 463)
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 454) (SEQ. ID NO: 464)
 40
            A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 455) (SEQ. ID NO: 465)
            A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 456) (SEQ. ID NO: 466)
            A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 457) (SEQ. ID NO: 467)
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 458) (SEQ. ID NO: 468)
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 459) (SEQ. ID NO: 469)
  45
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 460) (SEQ. ID NO: 470)
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 461) (SEQ. ID NO: 471)
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 462) (SEQ. ID NO: 472)
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 463) (SEQ. ID NO: 473)
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 464) (SEQ. ID NO: 474)
  50
         5'- A AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 465) (SEQ. ID NO: 475)
         5'- A AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 466) (SEQ. ID NO: 476)
            A AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 467) (SEQ. ID NO: 477)
          5'- A AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 468) (SEQ. ID NO: 478)
          5'- A AGC TGA GAT GGA GGG CGG CA-3' (FRAG 469) (SEQ. ID NO: 479)
  55
          5'- A AGC TGA GAT GGA GGG CGG C-3' (FRAG 470) (SEQ. ID NO: 480)
          5'- A AGC TGA GAT GGA GGG CGG -3' (FRAG 471) (SEQ. ID NO: 481)
          5'- A AGC TGA GAT GGA GGG CG -3' (FRAG 472) (SEQ. ID NO: 482)
          5'- A AGC TGA GAT GGA GGG C -3' (FRAG 473) (SEQ. ID NO: 483)
          5'- A AGC TGA GAT GGA GGG -3' (FRAG 474) (SEQ. ID NO: 484)
  60
          5'- A AGC TGA GAT GGA GG -3' (FRAG 475) (SEQ. ID NO: 485)
          5'- A AGC TGA GAT GGA G -3' (FRAG 476) (SEQ. ID NO: 486)
          5'- A AGC TGA GAT GGA -3' (FRAG 477) (SEQ. ID NO: 487)
          5'- A AGC TGA GAT GG -3' (FRAG 478) (SEQ. ID NO: 488)
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5'- A AGC TGA GAT G -3' (FRAG 479) (SEQ. ID NO; 489)
       5'- A AGC TGA GAT -3' (FRAG 480) (SEQ. ID NO: 490)
          AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 481) (SEQ. ID NO: 491)
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 482) (SEQ. ID NO: 492)
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 483) (SEQ. ID NO: 493)
5
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 484) (SEQ. ID NO: 494)
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 485) (SEQ. ID NO: 495)
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 486) (SEQ. ID NO: 496)
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 487) (SEQ. ID NO: 497)
       5'-
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 488) (SEQ. ID NO: 498)
10
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 489) (SEQ. ID NO: 499)
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 490) (SEQ. ID NO: 500)
           AGC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 491) (SEQ. ID NO: 501)
       5'-
            AGC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 492) (SEQ. ID NO: 502)
           AGC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 493) (SEQ. ID NO: 503)
15
            AGC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 494) (SEQ. ID NO: 504)
            AGC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 495) (SEQ. ID NO: 505)
            AGC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 496) (SEQ. ID NO: 506)
            AGC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 497) (SEQ. ID NO: 507)
            AGC TGA GAT GGA GGG CGG CAT G -3' (FRAG 498) (SEQ. ID NO: 508)
20
        51.
            AGC TGA GAT GGA GGG CGG CAT -3' (FRAG 499) (SEQ. ID NO: 509)
            AGC TGA GAT GGA GGG CGG CA-3' (FRAG 500) (SEQ. ID NO: 510)
            AGC TGA GAT GGA GGG CGG C-3' (FRAG 501) (SEQ. ID NO: 511)
             AGC TGA GAT GGA GGG CGG -3' (FRAG 502) (SEQ. ID NO: 512)
        5'-
            AGC TGA GAT GGA GGG CG -3' (FRAG 503) (SEQ. ID NO: 513)
 25
        5'-
             AGC TGA GAT GGA GGG C -3' (FRAG 504) (SEQ. ID NO: 514)
             AGC TGA GAT GGA GGG -3' (FRAG 505) (SEQ. ID NO: 515)
        5'-
             AGC TGA GAT GGA GG -3' (FRAG 506) (SEQ. ID NO: 516)
             AGC TGA GAT GGA G -3' (FRAG 507) (SEQ. ID NO: 517)
            AGC TGA GAT GGA -3' (FRAG 508) (SEQ. ID NO: 518)
 30
             AGC TGA GAT GG -3' (FRAG 509) (SEQ. ID NO: 519)
             AGC TGA GAT G -3' (FRAG 510) (SEQ. ID NO: 520)
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 511) (SEQ. ID NO: 521)
         5'-
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 512) (SEQ. ID NO: 522)
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 513) (SEQ. ID NO: 523)
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 514) (SEQ. ID NO: 524)
 35
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 515) (SEQ. ID NO: 525)
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 516) (SEQ. ID NO: 526)
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 517) (SEQ. ID NO: 527)
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 518) (SEQ. ID NO: 528)
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 519) (SEQ. ID NO: 529)
  40
             GC TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 520) (SEQ. ID NO: 530)
              GC TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 521) (SEQ. ID NO: 531)
              GC TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 522) (SEQ. ID NO: 532)
              GC TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 523) (SEQ. ID NO: 533)
              GC TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 524) (SEQ. ID NO: 534)
  45
              GC TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 525) (SEQ. ID NO: 535)
          5'-
              GC TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 526) (SEQ. ID NO: 536)
              GC TGA GAT GGA GGG CGG CAT GG -3' (FRAG 527) (SEQ. ID NO: 537)
              GC TGA GAT GGA GGG CGG CAT G -3' (FRAG 528) (SEQ. ID NO: 538)
          5'-
  50
              GC TGA GAT GGA GGG CGG CAT -3' (FRAG 529) (SEQ. ID NO: 539)
              GC TGA GAT GGA GGG CGG CA-3' (FRAG 530) (SEQ. ID NO: 540)
               GC TGA GAT GGA GGG CGG C-3' (FRAG 531) (SEQ. ID NO: 541)
          5'-
               GC TGA GAT GGA GGG CGG -3' (FRAG 532) (SEQ. ID NO: 542)
          5'-
               GC TGA GAT GGA GGG CG -3' (FRAG 533) (SEQ. ID NO: 543)
   55
               GC TGA GAT GGA GGG C -3' (FRAG 534) (SEQ. ID NO: 544)
               GC TGA GAT GGA GGG -3' (FRAG 535) (SEQ. ID NO: 545)
               GC TGA GAT GGA GG -3' (FRAG 536) (SEQ. ID NO: 546)
               GC TGA GAT GGA G -3' (FRAG 537) (SEQ. ID NO: 547)
               GC TGA GAT GGA -3' (FRAG 538) (SEQ. ID NO: 548)
   60
               GC TGA GAT GG -3' (FRAG 539) (SEQ. ID NO: 549)
               C TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 540) (SEQ. ID NO: 550)
               C TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 541) (SEQ. ID NO: 551)
           5.-
               C TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 542) (SEQ. ID NO: 552)
           5'-
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5'- C TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 543) (SEQ. ID NO: 553)
      5'- C TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 544) (SEQ. ID NO: 554)
           C TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 545) (SEQ. ID NO: 555)
          C TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 546) (SEQ. ID NO: 556)
           C TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 547) (SEQ. ID NO: 557)
5
          C TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 548) (SEQ. ID NO: 558)
           C TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 549) (SEQ. ID NO: 559)
           C TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 550) (SEQ. ID NO: 560)
           C TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 551) (SEQ. ID NO: 561)
           C TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 552) (SEQ. ID NO: 562)
10
           C TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 553) (SEQ. ID NO: 563)
           C TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 554) (SEQ. ID NO: 564)
           C TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 555) (SEQ. ID NO: 565)
           C TGA GAT GGA GGG CGG CAT GG -3' (FRAG 556) (SEQ. ID NO: 566)
           C TGA GAT GGA GGG CGG CAT G -3' (FRAG 557) (SEQ. ID NO: 567)
15
           C TGA GAT GGA GGG CGG CAT -3' (FRAG 558) (SEQ. ID NO: 568)
           C TGA GAT GGA GGG CGG CA-3' (FRAG 559) (SEQ. ID NO: 569)
            C TGA GAT GGA GGG CGG C-3' (FRAG 560) (SEQ. ID NO: 570)
            C TGA GAT GGA GGG CGG -3' (FRAG 561) (SEQ. ID NO: 571)
            C TGA GAT GGA GGG CG -3' (FRAG 562) (SEQ. ID NO: 572)
20
            C TGA GAT GGA GGG C -3' (FRAG 563) (SEQ. ID NO: 573)
            C TGA GAT GGA GGG -3' (FRAG 564) (SEQ. ID NO: 574)
            C TGA GAT GGA GG -3' (FRAG 565) (SEQ. ID NO: 575)
            C TGA GAT GGA G -3' (FRAG 566) (SEQ. ID NO: 576)
        5'-
            C TGA GAT GGA -3' (FRAG 567) (SEQ. ID NO: 577)
             TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 568) (SEQ. ID NO: 578)
25
             TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 569) (SEQ. ID NO: 579)
             TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 570) (SEQ. ID NO: 580)
        5'-
             TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 571) (SEQ. ID NO: 581)
             TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 572) (SEQ. ID NO: 582)
        5'-
 30
             TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 573) (SEQ. ID NO: 583)
             TGA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 574) (SEQ. ID NO: 584)
        5'-
             TGA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 575) (SEQ. ID NO: 585)
             TGA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 576) (SEQ. ID NO: 586)
        5'-
             TGA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 577) (SEQ. ID NO: 587)
 35
             TGA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 578) (SEQ. ID NO: 588)
             TGA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 579) (SEQ. ID NO: 589)
              TGA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 580) (SEQ. ID NO: 590)
              TGA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 581) (SEQ. ID NO: 591)
         5'-
              TGA GAT GGA GGG CGG CAT GGC G-3' (FRAG 582) (SEQ. ID NO: 592)
 40
              TGA GAT GGA GGG CGG CAT GGC -3' (FRAG 583) (SEQ. ID NO: 593)
              TGA GAT GGA GGG CGG CAT GG -3' (FRAG 584) (SEQ. ID NO: 594)
              TGA GAT GGA GGG CGG CAT G -3' (FRAG 585) (SEQ. ID NO: 595)
              TGA GAT GGA GGG CGG CAT -3' (FRAG 586) (SEQ. ID NO: 596)
              TGA GAT GGA GGG CGG CA-3' (FRAG 587) (SEQ. ID NO: 597)
  45
         5'-
              TGA GAT GGA GGG CGG C-3' (FRAG 588) (SEQ. ID NO: 598)
              TGA GAT GGA GGG CGG -3' (FRAG 589) (SEQ. ID NO: 599)
         5'-
              TGA GAT GGA GGG CG -3' (FRAG 590) (SEQ. ID NO: 600)
         5'-
              TGA GAT GGA GGG C -3' (FRAG 591) (SEQ. ID NO: 601)
              TGA GAT GGA GGG -3' (FRAG 592) (SEQ. ID NO: 602)
  50
              TGA GAT GGA GG -3' (FRAG 593) (SEQ. ID NO: 603)
          5'-
               TGA GAT GGA G -3' (FRAG 594) (SEQ. ID NO: 604)
               GA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 595) (SEQ. ID NO: 605)
               GA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 596) (SEQ. ID NO: 606)
          5'-
               GA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 597) (SEQ. ID NO: 607)
  55
               GA GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 598) (SEQ. ID NO: 608)
               GA GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 599) (SEQ. ID NO: 609)
          5'-
               GA GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3" (FRAG 600) (SEQ. ID NO: 610)
          5'-
               GA GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 601) (SEQ. ID NO: 611)
               GA GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 602) (SEQ. ID NO: 612)
  60
               GA GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 603) (SEQ. ID NO: 613)
               GA GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 604) (SEQ. ID NO: 614)
               GA GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 605) (SEQ. ID NO: 615)
               GA GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 606) (SEQ. ID NO: 616)
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GA GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 607) (SEQ. ID NO: 617)
            GA GAT GGA GGG CGG CAT GGC GG-3' (FRAG 608) (SEQ. ID NO: 618)
            GA GAT GGA GGG CGG CAT GGC G-3' (FRAG 609) (SEQ. ID NO: 619)
       5'-
            GA GAT GGA GGG CGG CAT GGC -3' (FRAG 610) (SEQ. ID NO: 620)
       5'-
            GA GAT GGA GGG CGG CAT GG -3' (FRAG 611) (SEQ. ID NO: 621)
 5
       5'-
            GA GAT GGA GGG CGG CAT G -3' (FRAG 612) (SEQ. ID NO: 622)
            GA GAT GGA GGG CGG CAT -3' (FRAG 613) (SEQ. ID NO: 623)
            GA GAT GGA GGG CGG CA-3' (FRAG 614) (SEQ. ID NO: 624)
       5'-
            GA GAT GGA GGG CGG C-3' (FRAG 615) (SEQ. ID NO: 625)
       51.
            GA GAT GGA GGG CGG -3' (FRAG 616) (SEQ. ID NO: 626)
10
            GA GAT GGA GGG CG -3' (FRAG 617) (SEQ. ID NO: 627)
            GA GAT GGA GGG C -3' (FRAG 618) (SEQ. ID NO: 628)
            GA GAT GGA GGG -3' (FRAG 619) (SEQ. ID NO: 629)
       5'-
             GA GAT GGA GG -3' (FRAG 620) (SEQ. ID NO: 630)
             A GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 621) (SEQ. ID NO: 631)
       51-
             A GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 622) (SEQ. ID NO: 632)
15
             A GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 623) (SEQ. ID NO: 633)
             A GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 624) (SEQ. ID NO: 634)
        51-
             A GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 625) (SEQ. ID NO: 635)
             A GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 626) (SEQ. ID NO: 636)
        5'-
20
             A GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 627) (SEQ. ID NO: 637)
             A GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 628) (SEQ. ID NO: 638)
             A GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 629) (SEQ. ID NO: 639)
             A GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 630) (SEQ. ID NO: 640)
        51.
             A GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 631) (SEQ. ID NO: 641)
25
             A GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 632) (SEQ. ID NO: 642)
             A GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 633) (SEQ. ID NO: 643)
             A GAT GGA GGG CGG CAT GGC GG-3' (FRAG 634) (SEQ. ID NO: 644)
             A GAT GGA GGG CGG CAT GGC G-3' (FRAG 635) (SEQ. ID NO: 645)
         5'-
              A GAT GGA GGG CGG CAT GGC -3' (FRAG 636) (SEQ. ID NO: 646)
 30
              A GAT GGA GGG CGG CAT GG -3' (FRAG 637) (SEQ. ID NO: 647)
              A GAT GGA GGG CGG CAT G -3' (FRAG 638) (SEQ. ID NO: 648)
         5'-
              A GAT GGA GGG CGG CAT -3' (FRAG 639) (SEQ. ID NO: 649)
              A GAT GGA GGG CGG CA-3' (FRAG 640) (SEQ. ID NO: 650)
         51-
              A GAT GGA GGG CGG C-3' (FRAG 641) (SEQ. ID NO: 651)
 35
              A GAT GGA GGG CGG -3' (FRAG 642) (SEQ. ID NO: 652)
              A GAT GGA GGG CG -3' (FRAG 643) (SEQ. ID NO: 653)
              A GAT GGA GGG C -3' (FRAG 644) (SEQ. ID NO: 654)
         5'-
              A GAT GGA GGG -3' (FRAG 645) (SEQ. ID NO: 655)
               GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 646) (SEQ. ID NO: 656)
         5'-
 40
               GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 647) (SEQ. ID NO: 657)
               GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 648) (SEQ. ID NO: 658)
         5'-
               GAT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 6) (SEQ. ID NO: 659)
         5'-
               GAT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 650) (SEQ. ID NO: 660)
          5'-
               GAT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 651) (SEQ. ID NO: 661)
 45
               GAT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 652) (SEQ. ID NO: 662)
               GAT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 653) (SEQ. ID NO: 663)
          5'-
               GAT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 654) (SEQ. ID NO: 664)
          5'-
               GAT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 655) (SEQ. ID NO: 665)
          5'-
               GAT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 656) (SEQ. ID NO: 666)
  50
          5'-
               GAT GGA GGG CGG CAT GGC GGG C-3' (FRAG 657) (SEQ. ID NO: 667)
          5'-
                GAT GGA GGG CGG CAT GGC GGG -3' (FRAG 658) (SEQ. ID NO: 668)
          5'-
                GAT GGA GGG CGG CAT GGC GG-3' (FRAG 659) (SEQ. ID NO: 669)
          5'-
                GAT GGA GGG CGG CAT GGC G-3' (FRAG 660) (SEQ. ID NO: 670)
          5'-
                GAT GGA GGG CGG CAT GGC -3' (FRAG 661) (SEQ. ID NO: 671)
  55
                GAT GGA GGG CGG CAT GG -3' (FRAG 662) (SEQ. ID NO: 672)
          5'-
                GAT GGA GGG CGG CAT G -3' (FRAG 663) (SEQ. ID NO: 673)
          5.-
                GAT GGA GGG CGG CAT -3' (FRAG 664) (SEQ. ID NO: 674)
          5'-
                GAT GGA GGG CGG CA-3' (FRAG 665) (SEQ. ID NO: 675)
          5'-
                GAT GGA GGG CGG C-3' (FRAG 666) (SEQ. ID NO: 676)
   60
                GAT GGA GGG CGG -3' (FRAG 667) (SEQ. ID NO: 677)
                GAT GGA GGG CG -3' (FRAG 668) (SEQ. ID NO: 678)
           5'-
                GAT GGA GGG C -3' (FRAG 669) (SEQ. ID NO: 679)
           5'-
                AT GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 670) (SEQ. ID NO: 680)
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AT GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 671) (SEQ. ID NO: 681)
             AT GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 672) (SEQ. ID NO: 682)
       5'-
             AT GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 673) (SEQ. ID NO: 683)
       5'-
             AT GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 674) (SEQ. ID NO: 684)
       5'-
             AT GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 675) (SEQ. ID NO: 685)
5
             AT GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 676) (SEQ. ID NO: 686)
       5'-
             AT GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 677) (SEQ. ID NO: 687)
       5'-
             AT GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 678) (SEQ. ID NO: 688)
             AT GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 679) (SEQ. ID NO: 689)
       5'-
             AT GGA GGG CGG CAT GGC GGG CA-3' (FRAG 680) (SEQ. ID NO: 690)
10
       51.
             AT GGA GGG CGG CAT GGC GGG C-3' (FRAG 681) (SEQ. ID NO: 691)
       5'-
             AT GGA GGG CGG CAT GGC GGG -3' (FRAG 682) (SEQ. ID NO: 692)
       5'-
             AT GGA GGG CGG CAT GGC GG-3' (FRAG 683) (SEQ. ID NO: 693)
             AT GGA GGG CGG CAT GGC G-3' (FRAG 684) (SEQ. ID NO: 694)
        5'-
             AT GGA GGG CGG CAT GGC -3' (FRAG 685) (SEQ. ID NO: 695)
15
             AT GGA GGG CGG CAT GG -3' (FRAG 686) (SEQ. ID NO: 696)
             AT GGA GGG CGG CAT G -3' (FRAG 687) (SEQ. ID NO: 697)
        5'-
             AT GGA GGG CGG CAT -3' (FRAG 688) (SEQ. ID NO: 698)
             AT GGA GGG CGG CA-3' (FRAG 689) (SEQ. ID NO: 699)
        51.
             AT GGA GGG CGG C-3' (FRAG 690) (SEQ. ID NO: 700)
20
        5'-
             AT GGA GGG CGG -3' (FRAG 691) (SEQ. ID NO: 701)
             AT GGA GGG CG -3' (FRAG 692) (SEQ. ID NO: 702)
        5'-
             T GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 693) (SEQ. ID NO: 703)
        5'-
             T GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 694) (SEQ. ID NO: 704)
        5'-
              T GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 695) (SEQ. ID NO: 705)
25
        5'-
             T GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 696) (SEQ. ID NO: 706)
        5'-
              T GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 697) (SEQ. ID NO: 707)
              T GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 698) (SEQ. ID NO: 708)
        5'-
              T GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 699) (SEQ. ID NO: 709)
        5'-
              T GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 700) (SEQ. ID NO: 710)
30
        51.
              T GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 701) (SEQ. ID NO: 711)
        5'-
              T GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 702) (SEQ. ID NO: 712)
        5'-
              T GGA GGG CGG CAT GGC GGG CA-3' (FRAG 703) (SEQ. ID NO: 713)
        5'-
              T GGA GGG CGG CAT GGC GGG C-3' (FRAG 704) (SEQ. ID NO: 714)
        5'-
              T GGA GGG CGG CAT GGC GGG -3' (FRAG 705) (SEQ. ID NO: 715)
 35
        5'-
              T GGA GGG CGG CAT GGC GG-3' (FRAG 706) (SEQ. ID NO: 716)
              T GGA GGG CGG CAT GGC G-3' (FRAG 707) (SEQ. ID NO: 717)
         5'-
              T GGA GGG CGG CAT GGC -3' (FRAG 708) (SEQ. ID NO: 718)
              T GGA GGG CGG CAT GG -3' (FRAG 709) (SEQ. ID NO: 719)
         51-
              T GGA GGG CGG CAT G -3' (FRAG 710) (SEQ. ID NO: 720)
 40
         51.
              T GGA GGG CGG CAT -3' (FRAG 711) (SEQ. ID NO: 721)
         5'-
              T GGA GGG CGG CA-3' (FRAG 712) (SEQ. ID NO: 722)
         5'-
              T GGA GGG CGG C-3' (FRAG 713) (SEQ. ID NO: 723)
              T GGA GGG CGG -3' (FRAG 714) (SEQ. ID NO: 724)
         5'-
               GGA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 715) (SEQ. ID NO: 725)
 45
         5'-
               GGA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 716) (SEQ. ID NO: 726)
         5'-
               GGA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 717) (SEQ. ID NO: 727)
         5'-
               GGA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 718) (SEQ. ID NO: 728)
                GGA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 719) (SEQ. ID NO: 729)
         5'~
                GGA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 720) (SEQ. ID NO: 730)
 50
         5'-
                GGA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 721) (SEQ. ID NO: 731)
         5'-
                GGA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 722) (SEQ. ID NO: 732)
         5'-
                GGA GGG CGG CAT GGC GGG CAC A-3' (FRAG 723) (SEQ. ID NO: 733)
         5'-
                GGA GGG CGG CAT GGC GGG CAC-3' (FRAG 724) (SEQ. ID NO: 734)
          5'-
                GGA GGG CGG CAT GGC GGG CA-3' (FRAG 725) (SEQ. ID NO: 735)
  55
          5'-
                GGA GGG CGG CAT GGC GGG C-3' (FRAG 726) (SEQ. ID NO: 736)
          5'-
                GGA GGG CGG CAT GGC GGG -3' (FRAG 727) (SEQ. ID NO: 737)
          5'-
                GGA GGG CGG CAT GGC GG-3' (FRAG 728) (SEQ. ID NO: 738)
                GGA GGG CGG CAT GGC G-3' (FRAG 729) (SEQ. ID NO: 739)
          5'-
                GGA GGG CGG CAT GGC -3' (FRAG 730) (SEQ, ID NO: 740)
  60
          5'-
                GGA GGG CGG CAT GG -3' (FRAG 731) (SEQ. ID NO: 741)
                GGA GGG CGG CAT G -3' (FRAG 732) (SEQ. ID NO: 742)
                GGA GGG CGG CAT -3' (FRAG 733) (SEQ. ID NO: 743)
                GGA GGG CGG CA-3' (FRAG 734) (SEQ. ID NO: 744)
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GGA GGG CGG C-3' (FRAG 735) (SEQ. ID NO: 745)
             GA GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 736) (SEQ. ID NO: 746)
             GA GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 737) (SEQ. 1D NO: 747)
       5'-
             GA GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 738) (SEQ. ID NO: 748)
       5'-
             GA GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 739) (SEQ. ID NO: 749)
5
       5'-
             GA GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 740) (SEQ. ID NO: 750)
       5'-
             GA GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 741) (SEQ. ID NO: 751)
       5'-
             GA GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 742) (SEQ. ID NO: 752)
       5'-
             GA GGG CGG CAT GGC GGG CAC AG-3' (FRAG 743) (SEQ. ID NO: 753)
       5'-
             GA GGG CGG CAT GGC GGG CAC A-3' (FRAG 744) (SEQ. ID NO: 754)
10
             GA GGG CGG CAT GGC GGG CAC-3' (FRAG 745) (SEQ. ID NO: 755)
       5'-
              GA GGG CGG CAT GGC GGG CA-3' (FRAG 746) (SEQ. ID NO: 756)
       5'-
              GA GGG CGG CAT GGC GGG C-3' (FRAG 747) (SEQ. ID NO: 757)
       5'-
              GA GGG CGG CAT GGC GGG -3' (FRAG 748) (SEQ. ID NO: 758)
       5'-
              GA GGG CGG CAT GGC GG-3' (FRAG 749) (SEQ. ID NO: 759)
15
        5'-
              GA GGG CGG CAT GGC G-3' (FRAG 750) (SEQ. ID NO: 760)
        5'-
              GA GGG CGG CAT GGC -3' (FRAG 751) (SEQ. ID NO: 761)
        5'-
              GA GGG CGG CAT GG -3' (FRAG 752) (SEQ. ID NO: 762)
        5'-
              GA GGG CGG CAT G -3' (FRAG 753) (SEQ. ID NO: 763)
        5'-
              GA GGG CGG CAT -3' (FRAG 754) (SEQ. ID NO: 764)
20
              GA GGG CGG CA-3' (FRAG 755) (SEQ. ID NO: 765)
        5'-
             A GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 756) (SEQ. ID NO: 766)
        5'-
              A GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 757) (SEQ. ID NO: 767)
        5'-
              A GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 758) (SEQ. ID NO: 768)
        5'-
              A GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 759) (SEQ. ID NO: 769)
25
              A GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 760) (SEQ. ID NO: 770)
              A GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 761) (SEQ. ID NO: 771)
              A GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 762) (SEQ. ID NO: 772)
        5'-
              A GGG CGG CAT GGC GGG CAC AG-3' (FRAG 763) (SEQ. ID NO: 773)
        5'-
              A GGG CGG CAT GGC GGG CAC A-3' (FRAG 764) (SEQ. ID NO: 774)
30
              A GGG CGG CAT GGC GGG CAC-3' (FRAG 765) (SEQ. ID NO: 775)
              A GGG CGG CAT GGC GGG CA-3' (FRAG 766) (SEQ. ID NO: 776)
        5'.
              A GGG CGG CAT GGC GGG C-3' (FRAG 767) (SEQ. ID NO: 777)
        5'-
              A GGG CGG CAT GGC GGG -3' (FRAG 768) (SEQ. ID NO: 778)
        5'-
              A GGG CGG CAT GGC GG-3' (FRAG 769) (SEQ. ID NO: 779)
 35
              A GGG CGG CAT GGC G-3' (FRAG 770) (SEQ. ID NO: 780)
              A GGG CGG CAT GGC -3' (FRAG 771) (SEQ. ID NO: 781)
              A GGG CGG CAT GG -3' (FRAG 772) (SEQ. ID NO: 782)
         5'-
               A GGG CGG CAT G -3' (FRAG 773) (SEQ. ID NO: 783)
         51-
               A GGG CGG CAT -3' (FRAG 774) (SEQ. ID NO: 784)
 40
               GGG CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 775) (SEQ. ID NO: 785)
               GGG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 776) (SEQ. ID NO: 786)
               GGG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 777) (SEQ. ID NO: 787)
         5'-
               GGG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 778) (SEQ. ID NO: 788)
         5'-
               GGG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 779) (SEQ. ID NO: 789)
 45
         5'-
               GGG CGG CAT GGC GGG CAC AGG C-3' (FRAG 780) (SEQ. ID NO: 790)
         5'-
                GGG CGG CAT GGC GGG CAC AGG -3' (FRAG 781) (SEQ. ID NO: 791)
                GGG CGG CAT GGC GGG CAC AG-3' (FRAG 782) (SEQ. ID NO: 792)
         5'-
                GGG CGG CAT GGC GGG CAC A-3' (FRAG 783) (SEQ. ID NO: 793)
         5'-
                GGG CGG CAT GGC GGG CAC-3' (FRAG 784) (SEQ. ID NO: 794)
  50
                GGG CGG CAT GGC GGG CA-3' (FRAG 785) (SEQ. ID NO: 795)
          5'-
                GGG CGG CAT GGC GGG C-3' (FRAG 786) (SEQ. ID NO: 796)
                GGG CGG CAT GGC GGG -3' (FRAG 787) (SEQ. ID NO: 797)
          5'-
                GGG CGG CAT GGC GG-3' (FRAG 788) (SEQ. ID NO: 798)
          5'-
                GGG CGG CAT GGC G-3' (FRAG 789) (SEQ. ID NO: 799)
  55
          5'-
                GGG CGG CAT GGC -3' (FRAG 790) (SEQ. ID NO: 800)
          5'-
                GGG CGG CAT GG -3' (FRAG 791) (SEQ. ID NO: 801)
          5'-
                GGG CGG CAT G -3' (FRAG 792) (SEQ. ID NO: 802)
          5'-
                GG CGG CAT GGC GGG CAC AG G CTG GGC-3' (FRAG 793) (SEQ. ID NO: 803)
          5'-
                GG CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 794) (SEQ. ID NO: 804)
  60
                GG CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 795) (SEQ. ID NO: 805)
                GG CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 796) (SEQ. ID NO: 806)
          5.
                GG CGG CAT GGC GGG CAC AGG CT-3' (FRAG 797) (SEQ. ID NO: 807)
          5'-
                GG CGG CAT GGC GGG CAC AGG C-3' (FRAG 798) (SEQ. ID NO: 808)
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GG CGG CAT GGC GGG CAC AGG -3' (FRAG 799) (SEQ. ID NO: 809)
       5'-
              GG CGG CAT GGC GGG CAC AG-3' (FRAG 800) (SEQ. ID NO: 810)
              GG CGG CAT GGC GGG CAC A-3' (FRAG 801) (SEQ. ID NO: 811)
       5'-
              GG CGG CAT GGC GGG CAC-3' (FRAG 802) (SEQ. ID NO: 812)
       5'-
              GG CGG CAT GGC GGG CA-3' (FRAG 803) (SEQ. ID NO: 813)
5
       5'-
              GG CGG CAT GGC GGG C-3' (FRAG 804) (SEQ. ID NO: 814)
       5'-
              GG CGG CAT GGC GGG -3' (FRAG 805) (SEQ. ID NO: 815)
       5'-
              GG.CGG CAT GGC GG-3' (FRAG 806) (SEQ. ID NO: 816)
        5'-
              GG CGG CAT GGC G-3' (FRAG 807) (SEQ. ID NO: 817)
        5'-
              GG CGG CAT GGC -3' (FRAG 808) (SEQ. ID NO: 818)
10
        ۲٠.
              GG CGG CAT GG -3' (FRAG 809) (SEQ. ID NO: 819)
        5'-
              G CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 810) (SEQ. ID NO: 820)
        5'-
              G CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 811) (SEQ. ID NO: 821)
        5'-
              G CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 812) (SEQ. ID NO: 822)
        51-
              G CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 813) (SEQ. ID NO: 823)
15
        5'-
              G CGG CAT GGC GGG CAC AGG CT-3' (FRAG 814) (SEQ. ID NO: 824)
        5'-
              G CGG CAT GGC GGG CAC AGG C-3' (FRAG 815) (SEQ. ID NO: 825)
               G CGG CAT GGC GGG CAC AGG -3' (FRAG 816) (SEQ. ID NO: 826)
        5'-
               G CGG CAT GGC GGG CAC AG-3' (FRAG 817) (SEQ. ID NO: 827)
        5'-
               G CGG CAT GGC GGG CAC A-3' (FRAG 818) (SEQ. ID NO: 828)
20
        5'-
               G CGG CAT GGC GGG CAC-3' (FRAG 819) (SEQ. ID NO: 829)
        5'-
               G CGG CAT GGC GGG CA-3' (FRAG 820) (SEQ. ID NO: 830)
        51.
               G CGG CAT GGC GGG C-3' (FRAG 821) (SEQ. ID NO: 831)
         5'-
               G CGG CAT GGC GGG -3' (FRAG 822) (SEQ. ID NO: 832)
         5'-
               G CGG CAT GGC GG-3' (FRAG 823) (SEQ. ID NO: 833)
25
        5'-
               G CGG CAT GGC G-3' (FRAG 824) (SEQ. ID NO: 834)
         5'-
               G CGG CAT GGC -3' (FRAG 825) (SEQ. ID NO: 835)
                CGG CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 826) (SEQ. ID NO: 836)
         5'-
                CGG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 827) (SEQ. ID NO: 837)
         5'-
                CGG CAT GGC GGG CAC AGG CTG G-3' (FRAG 828) (SEQ. ID NO: 838)
30
                CGG CAT GGC GGG CAC AGG CTG -3' (FRAG 829) (SEQ. ID NO: 839)
         5'-
                CGG CAT GGC GGG CAC AGG CT-3' (FRAG 830) (SEQ. ID NO: 840)
                CGG CAT GGC GGG CAC AGG C-3' (FRAG 831) (SEQ. ID NO: 841)
         5'-
                CGG CAT GGC GGG CAC AGG -3' (FRAG 832) (SEQ. ID NO: 842)
         5'-
                CGG CAT GGC GGG CAC AG-3' (FRAG 833) (SEQ. ID NO: 843)
 35
         5'-
                CGG CAT GGC GGG CAC A-3' (FRAG 834) (SEQ. ID NO: 844)
         5'•
                CGG CAT GGC GGG CAC-3' (FRAG 835) (SEQ. ID NO: 845)
          5'-
                 CGG CAT GGC GGG CA-3' (FRAG 836) (SEQ. ID NO: 846)
          5'-
                 CGG CAT GGC GGG C-3' (FRAG 837) (SEQ. ID NO: 847)
                 CGG CAT GGC GGG -3' (FRAG 838) (SEQ. ID NO: 848)
 40
          5'-
                 CGG CAT GGC GG-3' (FRAG 839) (SEQ. ID NO: 849)
          5'-
                 CGG CAT GGC G-3' (FRAG 840) (SEQ. ID NO: 850)
          5'-
                 GG CAT GGC GGG CAC AGG C TG GGC-3' (FRAG 841) (SEQ. ID NO: 851)
          5'-
                 GG CAT GGC GGG CAC AGG CTG GG-3' (FRAG 842) (SEQ, ID NO: 852)
          5'-
                 GG CAT GGC GGG CAC AGG CTG G-3' (FRAG 843) (SEQ. ID NO: 853)
  45
          5'-
                 GG CAT GGC GGG CAC AGG CTG -3' (FRAG 844) (SEQ. ID NO: 854)
GG CAT GGC GGG CAC AGG CT-3' (FRAG 845) (SEQ. ID NO: 855)
          5'-
          5'-
                 GG CAT GGC GGG CAC AGG C-3' (FRAG 846) (SEQ. ID NO: 856)
          5'-
                 GG CAT GGC GGG CAC AGG -3' (FRAG 847) (SEQ. ID NO: 857)
          5'-
                 GG CAT GGC GGG CAC AG-3' (FRAG 848) (SEQ. ID NO: 858)
  50
          5'-
                 GG CAT GGC GGG CAC A-3' (FRAG 849) (SEQ. ID NO: 859)
          5'-
                 GG CAT GGC GGG CAC-3' (FRAG 850) (SEQ. ID NO: 860)
          5'-
                  GG CAT GGC GGG CA-3' (FRAG 851) (SEQ. ID NO: 861)
          5'-
                  GG CAT GGC GGG C-3' (FRAG 852) (SEQ. ID NO: 862)
          5'-
                  GG CAT GGC GGG -3' (FRAG 853) (SEQ. ID NO: 863)
  55
          5'-
                  GG CAT GGC GG-3' (FRAG 854) (SEQ. ID NO: 864)
          5'-
                  G CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 855) (SEQ. ID NO: 865)
           5.
                  G CAT GGC GGG CAC AGG CTG GG-3' (FRAG 856) (SEQ. ID NO: 866)
           5'-
                  G CAT GGC GGG CAC AGG CTG G-3' (FRAG 857) (SEQ. ID NO: 867)
           5'-
                  G CAT GGC GGG CAC AGG CTG -3' (FRAG 858) (SEQ. ID NO: 868)
   60
           5'-
                  G CAT GGC GGG CAC AGG CT-3' (FRAG 859) (SEQ. ID NO: 869)
           5'.
                  G CAT GGC GGG CAC AGG C-3' (FRAG 860) (SEQ. ID NO: 870)
           5'-
                  G CAT GGC GGG CAC AGG -3' (FRAG 861) (SEQ. ID NO: 871)
           5'-
                  G CAT GGC GGG CAC AG-3' (FRAG 862) (SEQ. ID NO: 872)
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G CAT GGC GGG CAC A-3' (FRAG 863) (SEQ. ID NO: 873)
              G CAT GGC GGG CAC-3' (FRAG 864) (SEQ. ID NO: 874)
       5'-
              G CAT GGC GGG CA-3' (FRAG 865) (SEQ. ID NO: 875)
       5'-
              G CAT GGC GGG C-3' (FRAG 866) (SEQ. ID NO: 876)
       5'-
              G CAT GGC GGG -3' (FRAG 867) (SEQ. ID NO: 877)
5
       5'-
              CAT GGC GGG CAC AGG CTG GGC-3' (FRAG 868) (SEQ. ID NO: 878)
       5'-
              CAT GGC GGG CAC AGG CTG GG-3' (FRAG 869) (SEQ. ID NO: 879)
       5'-
              CAT GGC GGG CAC AGG CTG G-3' (FRAG 870) (SEQ. ID NO: 880)
       5'-
              CAT GGC GGG CAC AGG CTG -3' (FRAG 871) (SEQ. ID NO: 881)
       5'-
               CAT GGC GGG CAC AGG CT-3' (FRAG 872) (SEQ. ID NO: 882)
10
       5'-
               CAT GGC GGG CAC AGG C-3' (FRAG 873) (SEQ. ID NO: 883)
       5'-
               CAT GGC GGG CAC AGG -3' (FRAG 874) (SEQ. ID NO: 884)
       5'-
               CAT GGC GGG CAC AG-3' (FRAG 875) (SEQ. ID NO: 885)
       5'-
               CAT GGC GGG CAC A-3' (FRAG 876) (SEQ. ID NO: 886)
       5'-
               CAT GGC GGG CAC-3' (FRAG 877) (SEQ. ID NO: 887)
15
        5'-
               CAT GGC GGG CA-3' (FRAG 878) (SEQ. ID NO: 888)
        5'-
               CAT GGC GGG C-3' (FRAG 879) (SEQ. ID NO: 889)
        5 -
               AT GGC GGG CAC AGG CTG GGC-3' (FRAG 880) (SEQ. ID NO: 890)
        5'-
               AT GGC GGG CAC AGG CTG GG-3' (FRAG 881) (SEQ. ID NO: 891)
        5'-
               AT GGC GGG CAC AGG CTG G-3' (FRAG 882) (SEQ. ID NO: 892)
20
               AT GGC GGG CAC AGG CTG -3' (FRAG 883) (SEQ. ID NO: 893)
        5'-
               AT GGC GGG CAC AGG CT-3' (FRAG 884) (SEQ. ID NO: 894)
        5'-
               AT GGC GGG CAC AGG C-3' (FRAG 885) (SEQ. ID NO: 895)
        5'-
               AT GGC GGG CAC AGG -3' (FRAG 886) (SEQ. ID NO: 896)
        5'-
               AT GGC GGG CAC AG-3' (FRAG 887) (SEQ. ID NO: 897)
25
                AT GGC GGG CAC A-3' (FRAG 888) (SEQ. ID NO: 898)
        5'-
                AT GGC GGG CAC-3' (FRAG 889) (SEQ. ID NO: 899)
        5'-
                AT GGC GGG CA-3' (FRAG 890) (SEQ. ID NO: 900)
        5'-
                T GGC GGG CAC AGG CTG GGC-3' (FRAG 891) (SEQ. ID NO: 901)
        5'-
                T GGC GGG CAC AGG CTG GG-3' (FRAG 892) (SEQ. ID NO: 902)
30
                T GGC GGG CAC AGG CTG G-3' (FRAG 893) (SEQ. ID NO: 903)
                T GGC GGG CAC AGG CTG -3' (FRAG 894) (SEQ. ID NO: 904)
         5'-
                T GGC GGG CAC AGG CT-3' (FRAG 895) (SEQ. 1D NO: 905)
         5'-
                T GGC GGG CAC AGG C-3' (FRAG 896) (SEQ. ID NO: 906)
         5'-
                T GGC GGG CAC AGG -3' (FRAG 897) (SEQ. ID NO: 907)
 35
                T GGC GGG CAC AG-3' (FRAG 898) (SEQ. ID NO: 908)
                T GGC GGG CAC A-3' (FRAG 899) (SEQ. ID NO: 909)
                T GGC GGG CAC-3' (FRAG 900) (SEQ. ID NO: 910)
         51-
                 GGC GGG CAC AGG CTG GGC-3' (FRAG 901) (SEQ. ID NO: 911)
         5'-
                 GGC GGG CAC AGG CTG GG-3' (FRAG 902) (SEQ. ID NO: 912)
 40
                 GGC GGG CAC AGG CTG G-3' (FRAG 903) (SEQ. ID NO: 913)
         5'-
                 GGC GGG CAC AGG CTG -3' (FRAG 904) (SEQ. ID NO: 914)
                 GGC GGG CAC AGG CT-3' (FRAG 905) (SEQ. ID NO: 915)
         5'-
                 GGC GGG CAC AGG C-3' (FRAG 906) (SEQ. ID NO: 916)
         5'-
                 GGC GGG CAC AGG -3' (FRAG 907) (SEQ. ID NO: 917)
 45
                 GGC GGG CAC AG-3' (FRAG 908) (SEQ. ID NO: 918)
                 GGC GGG CAC A-3' (FRAG 909) (SEQ. ID NO: 919)
                 GC GGG CAC AGG CTG GGC-3' (FRAG 910) (SEQ. ID NO: 920)
         5'-
                 GC GGG CAC AGG CTG GG-3' (FRAG 911) (SEQ. ID NO: 921)
         5'-
                 GC GGG CAC AGG CTG G-3' (FRAG 912) (SEQ. ID NO: 922)
  50
                  GC GGG CAC AGG CTG -3' (FRAG 913) (SEQ. ID NO: 923)
          5'- GC GGG CAC AGG CT-3' (FRAG 914) (SEQ. ID NO: 924)
          5'- GC GGG CAC AGG C-3' (FRAG 915) (SEQ. ID NO: 925)
          5'- GC GGG CAC AGG -3' (FRAG 916) (SEQ. ID NO: 926)
          5'- GC GGG CAC AG-3' (FRAG 917) (SEQ. ID NO: 927)
  55
          5'- C GGG CAC AGG CTG GGC-3' (FRAG 918) (SEQ. ID NO: 928)
          5'- GGG CAC AGG CTG GG-3' (FRAG 919) (SEQ. ID NO: 929)
          5'- C GGG CAC AGG CTG G-3' (FRAG 920) (SEQ. ID NO: 930)
          5'- C GGG CAC AGG CTG -3' (FRAG 921) (SEQ. ID NO: 931)
          5'- C GGG CAC AGG CT-3' (FRAG 922) (SEQ. ID NO: 932)
  60
          5'- C GGG CAC AGG C-3' (FRAG 923) (SEQ. ID NO: 933)
          5'- C GGG CAC AGG -3' (FRAG 924) (SEQ. ID NO: 934)
          5'- GGG CAC AGG CTG GGC-3' (FRAG 925) (SEQ. ID NO: 935)
          5'- GGG CAC AGG CTG GG-3' (FRAG 926) (SEQ. ID NO: 936)
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5'- GGG CAC AGG CTG G-3' (FRAG 927) (SEQ. ID NO: 937)
       5'- GGG CAC AGG CTG -3' (FRAG 928) (SEQ. ID NO: 938)
       5'- GGG CAC AGG CT-3' (FRAG 929) (SEQ. ID NO: 939)
       5'- GGG CAC AGG C-3' (FRAG 930) (SEQ. ID NO: 940)
       5'- GG CAC AGG CTG GGC-3' (FRAG 931) (SEQ. ID NO: 941)
       5'- GG CAC AGG CTG GG-3' (FRAG 932) (SEQ. ID NO: 942)
       5'- GG CAC AGG CTG G-3' (FRAG 933) (SEQ. ID NO: 943)
                GG CAC AGG CTG -3' (FRAG 934) (SEQ. ID NO: 944)
                GG CAC AGG CT-3' (FRAG 935) (SEQ. ID NO: 945)
                G CAC AGG CTG GGC-3' (FRAG 936) (SEQ. ID NO: 946)
10
       5'-
                G CAC AGG CTG GG-3' (FRAG 937) (SEQ. ID NO: 947)
       5'-
                G CAC AGG CTG G-3' (FRAG 938) (SEQ. ID NO: 948)
       5'-
                G CAC AGG CTG -3' (FRAG 939) (SEQ. ID NO: 949)
                CAC AGG CTG GGC-3' (FRAG 940) (SEQ. ID NO: 950)
                 CAC AGG CTG GG-3' (FRAG 941) (SEQ. ID NO: 951)
15
        5'-CAC AGG CTG G-3' (FRAG 942) (SEQ. ID NO: 952)
        5'-AC AGG CTG GGC-3' (FRAG 943) (SEQ. ID NO: 953)
        5'-AC AGG CTG GG-3' (FRAG 944) (SEQ. ID NO: 954)
        5'-C AGG CTG GGC-3' (FRAG 945) (SEQ. ID NO: 955)
        5'-TTT TCC TTC CTT TGT CTC TCT TC (FRAG 946) (SEQ. ID NO: 956)
20
        5'-GCT CCC GGC TGC CTG (FRAG 947) (SEQ. ID NO: 957)
        5'-CTC GGC CGT GCG GCT CTG TCG CTC CCG GT (FRAG 948) (SEQ. ID NO: 958)
        5'-CCG CCG CCC TCC GGG GGG TC (FRAG 949) (SEQ. ID NO: 959)
        5'-TGC TGC CGT TGG CTG CCC (FRAG 950) (SEQ. ID NO: 960)
        5'-CTT CTG CGG GTC GCC GG (FRAG 951) (SEQ. ID NO: 961)
25
        5'-TGC TGG GCT TGT GGC (FRAG 952) (SEQ. ID NO: 962)
        5'-GGC CTC TCT TCT GGG (FRAG 953) (SEQ. ID NO: 963)
        5'-CCT GGT CCC TCC GT (FRAG 954) (SEQ. ID NO: 964)
        5'-GGT GGC TCC TCT GC (FRAG 955) (SEQ. ID NO: 965)
         5'-GCT TGG TCC TGG GGC TGC (FRAG 956) (SEQ. ID NO: 966)
 30
         5'-TGC TCT CCT CTC CTT (FRAG 957) (SEQ. ID NO: 967)
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Human Adenosine A2a Receptor Anti-sense Oligonucleotide Fragments

GCG GGT GGC CGT TG GGC CCG TGT TCC CCT GGG -GCC TGG GGC TCC CTT CTC TC GCC CTT CTT GCT GGG CCT C TGC TGC TGC TGC TGT GGC CCC C GTA CAC CGA GGA GCC CAT GAT GGG CAT GCC ACA GAC GAC AGG C GTB CBC 35 CGB GGB GCC CBT GBT GGG CBT GCC BCB GBC GBC BGG C-3' (FRAG. NO. 1665) (SEQ. ID NO:1678) 5'-CTG GGC CTC-3' (FRAG 1666) (SEQ. ID NO: 1679) 5'-GCC GCC CGC CTG-3' (FRAG 1667) (SEQ. ID NO: 1680) 5'-GC CCG CTC CCC GGC-3' (FRAG 1668) (SEQ. ID NO: 1681) 40 5'-CBCCGBGGBGCCC-3' (FRAG 1669) (SEQ. ID NO: 1682) 5'-TGC TTT TCT TTT CTG GGC CTC-3' (FRAG 958) (SEQ. ID NO: 968) 5'-TGT GGT CTG TTT TTT TCT G-3' (FRAG 959) (SEQ. ID NO: 969) 5'-GCC CTG CTG GGG CGC TCT CC-3' (FRAG 960) (SEQ. ID NO: 970) 5'-GCC GCC CGC CTG GCT CCC-3' (FRAG 961) (SEQ. ID NO: 971) 45 5'-GGB GCC CBT GBT GGG CBT GCC-3' (FRAG 962) (SEQ. ID NO: 972) 5'-GTG GTT CTT GCC CTC CTT TGG CTG-3' (FRAG 963) (SEQ. ID NO: 973) 5'-CCG TGC CCG CTC CCC GGC-3' (FRAG 964) (SEQ. ID NO: 974) 5'-CTC CTG GCG GGT GGC CGT TG-3' (FRAG 965) (SEQ. ID NO: 975) 5'-GGC CCG TGT TCC CCT GGG-3' (FRAG %6) (SEQ. ID NO: 976) 50

5'-GCC TGG GGC TCC CTT CTC TC-3' (FRAG 967) (SEQ. ID NO: 977) 5'-GCC CTT CTT GCT GGG CCT C-3' (FRAG 968) (SEQ. ID NO: 978)

55

5'-TGC TGC TGC TGG TGC TGT GGC CCC C-3' (FRAG 969) (SEQ. ID NO: 979)

5'-GTACACCGAGGAGCCCATGATGGGCATGCCACAGACGACAGGC-3' (FRAG 970) (SEQ. ID NO: 980) 5'-GTBCBCCGBGGGGCCCBTGBTGGGCBTGCCBCBGBCGBCGGC-3' (FRAG 971) (SEQ. ID NO: 981)

Human Adenosine A2b Receptor Anti-sense Oligonucleotide Fragments

5'-GGC GCC GTG CCG CGT CTT GGT GGC GGC GG GTT CGC GCC CGC GCG GGG CCC CTC CGG TCC GTT CGC GCC CGC

CGC CTC CGC CTG CCG CTT CTG GCT GGG CCC CGG GCG CCC CCT CCC CTC TTG CTC GGG TCC CCG TG ACA GCG CGT CCT GTG TCT CCA GCA GCA TGG CCG GGC CAG CTG GGC CCC BCB GCG CGT CCT GTG TCT CCB GCB GCB TGG CCG GGC CCT GTG TCT CCA GCA GCA TGG CCG GGC CAG CTG GGC CCC BCB GCG CBG CTG GGC CCC-3' (FRAG. NO: 1670) (SEQ. ID NO:1683)
5'-GCGCGTCCTG-3' (FRAG. NO: 1671) (SEQ. ID NO:1684)
5'-GCT GGG CCC CGG-3' (FRAG. NO: 1672) (SEQ. ID NO:1685)
5'-CGC GCC CGC G-3' (FRAG. NO: 1674) (SEQ. ID NO:1687)

- 5
- 5'- CGC GCC CGC G-3' (FRAG. NO: 16/4) (SEQ. ID NO:1867)
 5'-GGC GCC GTG CCG CGT CTT GGT GGC GGC GG-3' (FRAG 972) (SEQ. ID NO: 982)
 5'-GTT CGC GCC CGC GCG GGG CCC CTC CGG TCC-3' (FRAG 973) (SEQ. ID NO: 983)
 5'-GTT CGC GCC CGC GCG GGG CCC CTC CGG TCC-3' (FRAG 974) (SEQ. ID NO: 984) 10
- 5'-CGG GTC GGG GCC CCC CGC GGC C-3' (FRAG 975) (SEQ. ID NO: 985)
 - 5'-GCC TCG GGG CTG GGG CGC TGG TGG CCG GG-3' (FRAG 976) (SEQ. ID NO: 986)
 - 5'-CCG CGC CTC CGC CTG CCG CTT CTG-3' (FRAG 977) (SEQ. ID NO: 987)
 - 5'-GCT GGG CCC CGG GCG CCC CCT-3' (FRAG 978) (SEQ. ID NO: 988)
- 5'-CCC CTC TTG CTC GGG TCC CCG TG-3' (FRAG 979) (SEQ. ID NO: 989) 15
 - 5'-ACAGCGCGTCCTGTGTCTCCAGCAGCATGGCCGGGCCAGCTGGGCCCC-3' (FRAG 980) (SEQ. ID NO: 990)
 - 5'-BCBGCGCGTCCTGTGTCTCCBGCBGCBTGGCCGGGCCBGCTGGGCCCC-3' (FRAG 981) (SEQ. ID NO: 991)

Human Adenosine A3 Receptor Anti-sense Oligonucleotide Fragments

- 5'-ACA GAG CAG TGC TGT TGT TGG GCA TCT TGC CTT CCC AGG G BCB GBG CB TGC TGT TGT TGG GCB TCT TGC CTT 20 NO:1675) (SEQ. ID NO:1688)
 - 5'-GBG CB TGC-3' (FRAG. NO:1676) (SEQ. ID NO:1689)
- 5'-TTG TTG GGC-3' (FRAG. NO:1677) (SEQ. ID NO:1690)
- 5'-TGC CTT CCC BGG G-3' (FRAG. NO:1678) (SEQ. ID NO:1691) 25
 - 5'-GTT GTT GGG CAT CTT GCC-3' (FRAG. NO:1679) (SEQ ID NO:3)
 - 5'-GTG GGC CTA GCT CTC GCC-3' (GRAG. NO:1680) (SEQ ID NO:5)
 - 5'-ACA GAG CA TGC TGT TGT TGG GCA TCT TGC CTT CCC AGG G-3' (FRAG 982) (SEQ. ID NO: 992)
 - 5'-BCB GBG CB TGC TGT TGT TGG GCB TCT TGC CTT CCC BGG G-3' (FRAG 983) (SEQ. ID NO: 993)
- 5'-CCC TTT TCT GGT GGG GTG-3' (FRAG 984) (SEQ. ID NO: 994) 30
 - 5'-GTG CTG TTG TTG GGC-3' (FRAG 985) (SEQ. ID NO: 995)
 - 5'-TTT CTT CTG TTC CC-3' (FRAG 986) (SEQ. ID NO: 996)
 - 5'-CCC TTT TCT GGT GGG GTG-3' (FRAG 987) (SEQ. ID NO: 997)
 - 5'-GTG CTG TTG TTG GGC-3' (FRAG 988) (SEQ. ID NO: 998)
- 5'-TTT CTT CTG TTC CC-3' (FRAG 989) (SEQ. ID NO: 999) 35

Human IgE Receptor β Anti-sense Oligonucleotide Fragments

AAC CGA GCT GT BTT TGC TCT CCT BTT BCT TTC TGT GTC CBT TTT TTC BTT BBC CGB GCT GT-3' (FRAG. NO:1681) (SEQ. ID NO:1692)

- 5'-CCC CTG GG-3' (FRAG. NO:1682) (SEQ. ID NO:1693) 40
 - 5'-GCTCTCCTBTT-3' (FRAG. NO:1683) (SEQ. ID NO:1694)
 - 5'-CBTTBBCCGBGCTG-3' (FRAG. NO:1684) (SEQ. ID NO:1695)
 - 5'-TTT CCC CTG GGT CTT CC-3' (FRAG 990) (SEQ. ID NO: 1000)
 - 5'-CTC CTG CTC TTT TTT C-3' (FRAG 991) (SEQ. ID NO: 1001)
- ATTTGCTCTCCTATTACTTTCTGTGTCCATTTTTTCATTAACCGAGCTGT (FRAG 992) (SEQ. ID NO: 1002) 45 BTTTGCTCTCCTBTTBCTTTCTGTGTCCBTTTTTCBTTBBCCGBGCTGT (FRAG 993) (SEQ. ID NO: 1003)

Human Fc-ξ Receptor CD23 Antigen (IgE Receptor) Antisense Oligonucleotide Fragments

- 5'-GCC TGT GTC TGT CCT CCT GCT TCG TTC CTC TCG TTC CTG CTT GGT GCC GTT GCC G GTC CTG CTC CGG GCT 50 TCT GGG CCT GGC TGT GGC CGT GGT TGG GGG TCT TC GCT GCC TCC GTT TGG GTG GC TCT CTG AAT ATT GAC CTT CCT CCA TGG CGG TCC TGC TTG GAT TCT CCC GA TCT CTG BBT BTT GBC CTT CCT CCB TGG CGG TCC TGC TTG GBT TCT CCC GB-3' (FRAG 1685) (SEQ. ID NO:1696)
- 5'-GT CCT CCT-3' (FRAG 1686) (SEQ. ID NO: 1697)
- 5'-TGT GTC TGT CCT CC-3' (FRAG 1687) (SEQ. ID NO: 1698) 55
 - 5'-GTG GCC CTG GC-3' (FRAG 1688) (SEQ. ID NO: 1699)
 - 5'-CGT GGT TGG GG-3' (FRAG 1689) (SEQ. ID NO: 1700) 5'-TCT CTG BBT BTT GBC C-3' (FRAG1690) (SEQ. ID NO:1701)
 - 5'-GCC TGT GTC TGT CCT CCT-3' (FRAG 994) (SEQ. ID NO: 1004)

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5'-GCT TCG TTC CTC TCG TTC-3' (FRAG 995) (SEQ. ID NO:1005)
5'-CTG CTT GGT GCC CTT GCC G-3' (FRAG 996) (SEQ. ID NO: 1006)
5'-GTC CTG CTC CGG GCT GTG G-3' (FRAG 997) (SEQ. ID NO: 1007)
5'-GTC GTG GCC CTG GCT CCG GCT GGT GGG CTC CCC TGG-3' (FRAG 998) (SEQ. ID NO: 1008)
5'-CCT TCG CTG GCT GGC GGC GTG C-3' (FRAG 999) (SEQ. ID NO: 1009)
5'-GGG TCT TGC TCT GGG CCT GGC TGT-3' (FRAG 1000) (SEQ. ID NO: 1010)
5'-GGC CGT GGT TGG GGG TCT TC-3' (FRAG 1001) (SEQ. ID NO: 1011)
5'-GCT GCC TCC GTT TGG GTG GC (FRAG 1002) (SEQ. ID NO: 1012)
5'-TCT CTG AAT ATT GAC CTT CCT CCA TGG CGG TCC TGC TTG GAT TCT CCC GA (FRAG 1003) (SEQ. ID NO: 1013)
5'-TCT CTG BBT BTT GBC CTT CCT CCB TGG CGG TCC TGC TTG GBT TCT CCC GB (FRAG 1004) (SEQ. ID NO: 1014)
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Human IgE Receptor α Subunit Anti-sense Oligonucleotide Fragments

5'- GCC TIT CCT GGT TCT CTT GTT GTT TTT GGG GTT TGG CTT ACA GTA GAG TAG GGG ATT CCA TGG CAG GAG CCA TCT TCA TGG ACT CC TTC AAG GAG ACC TTA GGT TTC TGA GGG ACT GCT AAC ACG CCA TCT GGA GC BCB GTB GBG TBG GGG BTT CCB TGG CBG GBG CCB TCT TCT TCB TGG BCT CC TTC BBG GBG BCC TTB GGT TTC TGB GGG BCT GCT BBC BCG CCB TCT GGB GC GTT GTT TTT GGG GTT TGG CTT GCC TTT CCT GGT TCT CTT BCB GTB GBG TBG GGG 15 BTT CCB TGG CBG GBG CCB TCT TCT TCB TGG BCT CC TTC BBG GBG BCC TTB GGT TTC TGB GGG BCT GCT BBC BCG CCB TCT GGB GC-3

(FRAG. NO: 1691) (SEQ. ID NO:1702)

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5'- TGG BCT CC -3' (FRAG. NO: 1692) (SEQ. ID NO:1703)

5'-CCB TCT GGB-3' (FRAG. NO: 1693) (SEQ. ID NO:1704) 20

5'-CT GCT BBC BCG-3' (FRAG. NO: 1694) (SEQ. ID NO:1705)

5'-GTT TTT GGG GTT TG-3' (FRAG. NO: 1695) (SEQ. ID NO:1706)

5'-GCC TIT CCT GGT TCT CTT GTT GTT TTT GGG GTT TGG CTT-3' (FRAG. NO:1005) (SEQ. ID NO:1015)

5'-ACAGTAGAGTAGGGGATTCCATGGCAGGAGCCATCTTCTTCATGGACTCC-3' (FRAG. NO:1006) (SEQ. ID NO:1016)

5'-TTC AAG GAG ACC TTA GGT TTC TGA GGG ACT GCT AAC ACG CCA TCT GGA GC-3' (FRAG. NO:1007) (SEQ. ID 25 NO-1017)

5'-BCB GTB GBG TBG GGG BTT CCB TGG CBG GBG CCB TCT TCT TCB TGG BCT CC TTC BBG GBG BCC TTB GGT TTC TGB GGG-3' (FRAG. NO:1008) (SEQ. ID NO:1018)

5'-BCT GCT BBC BCG CCB TCT GGB GC-3' (FRAG, NO:1009) (SEQ. ID NO:1019)

5'-GTT GTT TTT GGG GTT TGG CTT-3' (FRAG. NO:1010) (SEQ. ID NO:1020) 30

5'-GCC TTT CCT GGT TCT CTT-3' (FRAG. NO:1011) (SEQ. ID NO:1021) 5'-BCBGTBGBGGGGBTTCCBTGGCBGGBGCCBTCTTCTTCBTGGBCTCC-3' (FRAG. NO:1012) (SEQ. ID NO:1022)

5'-TTC BBG GBG BCC TTB GGT TTC TGB GGG BCT GCT BBC BCG CCB TCT GGB GC-3' (FRAG. NO:1013) (SEQ. ID NO > 1023)

Human IgE Receptor (Fc Epsilon R) Anti-sense Oligonucleotide Fragments 35

5'-GCC TGT GTC TGT CCT CCT GCT TCG TTC CTC TCG TTC CTG GCT GCC G GTC CTG CTC CGG GCT CCC GGB CCG BCB GGC CGT GGT TGG GGG TCT TC GCT GCC TCC GTT TGG GTG GC GAT CTC TGA ATA TTGA CCT TCC ATG GCG GTC CTG GTT GGA GBT CTC TGB BTB TTGB CCT TCC BTG GCG GTC CTG CTT GGB-3' (FRAG: 1696) (SEQ. ID NO:1707)

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5'-TCG TTC CTC TCG-3' (FRAG: 1697) (SEQ. ID NO:1708)
5'-BGB BCG BGB C-3' (FRAG: 1698) (SEQ. ID NO:1709)
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5'-TGB BTB TTGB-3' (FRAG: 1699) (SEQ. ID NO:1710)

5'-GCC TGT GTC TGT CCT CCT-3' (FRAG. NO:1014) (SEQ. ID NO:1024)

5'-GCT TCG TTC CTC TCG TTC-3' (FRAG. NO:1015)(SEQ. ID NO:1025)

5'-CTG CTT GGT GCC CTT GCC G-3' (FRAG. NO:1016)(SEQ. ID NO:1026)

5'-GTC CTG CTC CGG GCT GTG G-3' (FRAG. NO:1017)(SEQ. ID NO:1027) 5'-GTC CTC GCC CTG GCT CCG GCT GGT GGG CTC CCC TGG-3' (FRAG. NO:1018) (SEQ. ID NO:1028)

5'-CCT TCG CTG GCT GGC GGC GTG C-3' (FRAG. NO:1019) (SEQ. ID NO:1029)

5'-CCC BGB BCG BGB CCC GGB CCG BCB-3' (FRAG. NO:1020) (SEQ. ID NO:1030) 50

5'-GGC CGT GGT TGG GGG TCT TC-3' (FRAG. NO:1021) (SEQ. ID NO:1031)

5'-GCT GCC TCC GTT TGG GTG GC-3' (FRAG. NO:1022) (SEQ. ID NO:1032)

5'-GBT CTC TGB BTB TTGB CCT TCC BTG GCG GTC CTG CTT GGB-3' (FRAG. NO:1023) (SEQ. ID NO:1033)

Human Histidine Decarboxylase Anti-sense Oligonucleotide Fragments

55 CTC BGG CTC CBT CBT CTC CCT TGG GC-3' (FRAG. NO:1700) (SEQ. ID NO:1711)

(FRAG. NO:1701) (SEQ. ID NO: 1712) 5'-GGC TCT GGC (FRAG. NO:1702) (SEQ. ID NO: 1713) 5'-CCC TTG G (FRAG. NO:1703) (SEQ. ID NO: 1714) 5'- TT TGT TCT TCC 5'- TCT CCC TTG GGC TCT GGC TCC TTC TC-3' (FRAG. NO:1024) (SEQ. ID NO: 1034) 5'- TCT CTC TCC CTC TCT CTC TGT -3' (FRAG. NO:1025) (SEQ. ID NO:1035) 5'- TTT TGT TCT TCC TTG CTG CC-3' (FRAG. NO:1027) (SEQ. ID NO:1037) 5'- GCC CCG CTG CTT GTC T TC CTC G-3' (FRAG. NO:1028) (SEQ. ID NO:1038) 5'-CTC TGT CCC TCT CTC TCT GTB CTC CTC BGG CTC CBT CBT CTC CCT TGG GC (FRAG. NO:1029) (SEQ. ID NO:1039)

Human Beta Tryptase Anti-sense Oligonucleotide Fragments 10

CGC GGC TCG CCB GGB CGG GCB GCB GCB GCB GCB GBT TCB GCB TCC TGG-3' (FRAG. NO:1704) (SEQ. ID NO: 1715)

5'- GCT CCT GGG GGC CT-3' (FRAG. NO:1705) (SEQ. ID NO: 1716)
5'-CGT BGG CGC-3' (FRAG. NO:1706) (SEQ. ID NO: 1717)
5'-T GGC CTG GGG-3' (FRAG. NO:1707) (SEQ. ID NO: 1718) 15 5'-CTT GCT CCT GGG GGC CTC CTG-3' (FRAG. NO:1030) (SEQ. ID NO:1040) 5'-GTC CCT CCG GGT GTT CCC GGC-3' (FRAG. NO:1031) (SEQ. ID NO:1041)

5'-GGG CCT GGC CTG GGG CBG GGG CCG CGT BGG CGC GGC TCG CCB GGB CGG GCB GCB GCB GCB GCB GBT TCB

GCB TCC TGG-3' (FRAG. NO:1032) (SEQ. ID NO:1042) 20

Human Tryptase-I Anti-sense Oligonucleotide Fragments

5'-CTT GCT CCT GGG GGC CTC CTG GTC CCT CTG GCT G TT CCC GGC CCT GGB CTG GGG CBG GGG CCG CGT BGG CGC GGC TCG CCB GGB CGG GCB GCC GCB GCB GCB GCC TCB GCC TCG CCB CGG BBT TCC-3' (FRAG. NO: 1708) (SEQ. ID NO:1719)

5'-CT CCT GGG GGC CTC CTG-3' (FRAG. NO:1709) (SEQ. ID NO:1720) 25 5'-B TCC TGG CCB CGG BBT TCC -3' (FRAG. NO:1710) (SEQ. ID NO:1721) 5'-GTC CCT C-3' (FRAG. NO:1711) (SEQ. ID NO:1722) 5'-CTT GCT CCT GGG GGC CTC CTG-3' (FRAG. NO:1033) (SEQ. ID NO:1043) 5'-GTC CCT CTG GCT G TT CCC GGC-3' (FRAG. NO:1034) (SEQ. ID NO:1044)

30 TCC TGG CCB CGG BBT TCC -3' (FRAG. NO:1035) (SEQ. ID NO:1045)

Human Prostaglandin D Synthase Anti-sense Oligonucleotide Fragments

GGG GCC CTT GCT GCC CTG GCT GT GCC CTG GGG GTC TGG GTT CGG CTG T CCC CBG CBG GBC CBG TCC CBT CCB CBG CGT GTG BTG BGT BGC CBT TCT CCT GCB GCC GBG-3' (FRAG. NO:1712) (SEQ. ID NO:1723)

35 5'-T TCT CCT GCB GCC GBG -3' (FRAG, NO:1713) (SEQ. ID NO:1724) 5'-CTT GCT GCC CTG GCT GT-3' (FRAG. NO:1714) (SEQ. ID NO:1725)

5'- TCT TCT CCT GG-3' (FRAG. NO:1715) (SEQ. ID NO:1726)

5'-GGT GTG CGG GGC CTG GTG CC-3' (FRAG. NO:1036) (SEQ. ID NO:1046)

5'-CCT GGG CCT CGG GTG CTG CCT GT-3' (FRAG. NO:1037) (SEQ. ID NO:1047) 40

5'-GCG CTG CCT TCT TCT CCT GG-3' (FRAG. NO:1038) (SEQ. ID NO:1048) 5'-GTC CTC GCC GGG GCC CTT GCT GCC CTG GCT GT-3' (FRAG. NO:1039) (SEQ. ID NO:1049)

5'-GCC CTG GGG GTC TGG GTT CGG CTG T-3' (FRAG. NO:1040) (SEQ. ID NO:1050)

5'-CCC CBG CBG GBC CBG TCC CBT CCB CBG CGT GTG BTG BGT BGC CBT TCT CCT GCB GCC GBG -3'

(FRAG. NO:1041) (SEQ. ID NO:1051) 45

Human Cyclooxygenase-2 Anti-sense Oligonucleotide Fragments

5'-GGG CGC GGG CGB GCB TCG C TTT GGG CTT TTC TCC TTT GGT T TGB GCG CCB GGB CCG CGC BCB GCB GCB GGG CGC GGG CGB GCB TCG CBG CGG CGG GCB GGG-3' (FRAG. NO: 1716) (SEQ. ID NO:1727)

5'-G GCB GGG -3' (FRAG. NO: 1717) (SEQ. ID NO: 1728) 5'-TCC TTT GGT T-3' (FRAG. NO:1718) (SEQ. ID NO:1729) 50

5'- GGG CGC GGG CGB GCB TCG C-3' (FRAG. NO:1042) (SEQ. ID NO:1052) 5'- TTT GGG CTT TTC TCC TTT GGT T-3' (FRAG. NO:1043) (SEQ. ID NO:1053)

5'-TGB GCG CCB GGB CCG CGC BCB GCB GCB GGG CGC GGG CGB GCB TCG CBG CGG CGG GCB GGG -3'

(FRAG. NO:1044) (SEQ. ID NO:1054)

Human Eosinophil Cationic Protein Anti-sense Oligonucleotide Fragments 55

5'-CCT CCT TCC TGG TCT GTC TGC CBG BCB BBT TTG GGB BGT GBB CBG TTT TGG BBC CBT GTT TCC CBG TCT CTG BGC TGT GGC-3' (FRAG. NO: 1719) (SEQ. ID NO: 1730) 5'-TTC TCC TTT GGT T-3' (FRAG. NO:1720) (SEQ. ID NO: 1731) 5'-T TTC TCC TTT GGT T-3' (FRAG. NO:1721) (SEQ. ID NO:1732)

5'- GGG CGC GGG CGB GCB TCG C-3' (FRAG. NO:1042) (SEQ. ID NO:1052) 5'- TTT GGG CTT TTC TCC TTT GGT T-3' (FRAG. NO:1043) (SEQ. ID NO:1053) (FRAG. NO:1044) (SEQ. ID NO:1054)

Human Eosinophil Derived Neurotoxin Anti-sense Oligonucleotide Fragments

TCC C TGT TTG CTG GTG TCT GCG C 5'- CCC CBB CBG BBG CBG BCB BBT TTG GGB BGT GBB CBG TTT TGG BBC CBT 10 GTT TCC TGT-3'

(FRAG. NO: 1722) (SEQ. ID NO: 1733)

5'-TTC CTG T-3' (FRAG. NO:1723) (SEQ. ID NO: 1734)

- 5'-CTC TTT CTG CT-3' (FRAG. NO: 1724) (SEQ. ID NO:1735) 15
 - 5'-CCC CTT CTG TCC C-3' (FRAG. NO:1725) (SEQ. ID NO: 1736)
 - 5'- GCC CTG CTG CTC TTT CTG CT-3' (FRAG. NO:1047) (SEQ. ID NO:1057)
 - 5'- TCC CTT GGT GGG TTG GGC C-3' (FRAG. NO:1048) (SEQ. ID NO:1058) 5'- GCT GGT TGT TCT GGG GTT C-3' (FRAG. NO:1049) (SEQ. ID NO:1059)
- 5'-TTG CTG CCC CTT CTG TCC C-3' (FRAG. NO:1050) (SEQ. ID NO:1060)
- 20 5'- TGT TTG CTG GTG TCT GCG C -3' (FRAG. NO:1051) (SEQ. ID NO:1061)
 - 5'- CCC CBB CBG BBG CBG BCB BBT TTG GGB BGT GBB CBG TTT TGG BBC CBT GTT TCC TGT-3' (FRAG. NO:1052) (SEQ. ID NO:1062)

Human Eosinophil Peroxidase Anti-sense Oligonucleotide Fragments

- 25 TTT GGG GGC CGG CGT TGT CTT G GTT TGG GGG TTT CCG TTG GGG TTC TCC TGG CCC GGG CCT TGC CC TCC BGT GGT GCG GTB CTT GTC GCT GCB GCG CTC GGC CTG GTC CCG GBG BGC -3' (FRAG. NO: 1726) (SEQ. ID
- 5'-TC GGC CTG GTC CCG G-3' (FRAG. NO: 1727) (SEQ. ID NO:1738) 30 5'-TGG GGG TTT CCG TTG-3' (FRAG. NO: 1728) (SEQ. ID NO: 1739) 5'-TG GTC CCG GBG BGC -3' (FRAG. NO: 1729) (SEQ. ID NO: 1740) 5'-GCG CTC GGC CTG GTC CCG G-3' (FRAG. NO:1053) (SEQ. ID NO:1063)
 - 5'-GGG TCT CCT CTT GTT GTT GC-3' (FRAG. NO:1054) (SEQ. ID NO:1064)
- 5'- TTG CGC CTC CTG CTG GGG GT CC-3' (FRAG. NO:1055) (SEQ. ID NO:1065) 35
 - 5'-CTC TGT TCT TGT TTT GGG GGC-3' (FRAG. NO:1056) (SEQ. ID NO:1066)
 - 5'-GGG CCC GGC CGT TGT CTT G-3' (FRAG. NO:1057) (SEQ. ID NO:1067)
 - 5'-GTT TGG GGG TTT CCG TTG-3' (FRAG. NO:1058) (SEQ. ID NO:1068)
 - 5'-GGG TTC TCC TGG CCC GGG CCT TGC CC-3' (FRAG. NO:1059)(SEQ. ID NO:1069) 5'-GGC CGT GGT CCC GGC TTC GTT GC-3' (FRAG. NO:1060) (SEQ. ID NO:1070)
- 40 5'-CCT GTC TCC GTC TCG GCT CTT CTG-3' (FRAG. NO:1061) (SEQ. ID NO:1071)
 - 5'-GGG CCT TGC GCT GTC TTT GGT G-3' (FRAG. NO:1062) (SEQ. ID NO:1072)
 - 5'-GCB CCG TCC BGT GBT GGT GCG GTB CTT GTC GCT GCB GCG CTC GGC CTG GTC CCG GBG BGC -3' (FRAG. NO:1063) (SEQ. 1D NO:1073)

Human Intercellular Adhesion Molecule-1 (ICAM-1) 45 Anti-sense Oligonucleotide Fragments

GTG TCT CCT GGC TCT GGT TCC CC GCT GCG CCC GTT GTC CTC TGG GGT GGC CTT C GCT CCC GGG TCT GGT TCT TGT CGG BGC CTC CCC GGG GCB GGB TGB CTT TTG BGG GGG BCB CBG BTG TCT GGG CBT TGC CBG GTC CTG GGB BCB GBG 50 CCC CGB GCB GGB CCB GGB GTG CGG GCB GCG CGG GCC GGG GGC TGC TGG GBG CCB TBG CGB GGC TGB G-3 (FRAG. NO: 1730)

(SEQ. ID NO: 1741)

- 5'-GGG GGC TGC TGG G-3' (FRAG. NO: 1731) (SEQ. ID NO:1742)
- 5'-T GTC CTC CGG CGT CCC-3' (FRAG. NO:1732) (SEQ. ID NO:1743) 55
 - 5'-G CCB TBG CGB GGC TGB G-3' (FRAG. NO: 1733) (SEQ. ID NO: 1744) 5'-CTC TGG GGT GGC CTT C-3' (FRAG. NO:1734) (SEQ. ID NO:1745)
 - 5'-GCG CGG GCC GGG GGC TGC TGG G-3' (FRAG. NO:1064) (SEQ. ID NO:1074)

PCT/US98/19419

WO 99/13886

5'-GGT TGG CCC GGG GTG CCC C-3' (FRAG. NO:1065) (SEQ. ID NO:1075) 5'-GCC GCT GGG TGC CCT CGT CCT CTG CGG TC-3' (FRAG, NO:1066) (SEQ. ID NO:1076) 5'-GTG TCT CCT GGC TCT GGT TCC CC-3' (FRAG. NO:1067) (SEQ. ID NO:1077) 5'-GCT GCG CCC GTT GTC CTC TGG GGT GGC CTT C-3' (FRAG. NO:1068) (SEQ. ID NO:1078) 5'-GCT CCC GGG TCT GGT TCT TGT GT-3' (FRAG. NO:1069) (SEQ. ID NO:1079) 5'-TGG GGG TCC CTT TTT GGG CCT GTT GT-3' (FRAG. NO:1070) (SEQ. ID NO:1080) 5'-GGC GTG GCT TGT GTG TTC GGT TTC-3' (FRAG. NO:1071) (SEQ. ID NO:1081) 5'-TGC CCT GTC CTC CGG CGT CCC-3' (FRAG. NO:1072) (SEQ. ID NO:1082) 5'- CGG BGC CTC CCC GGG GCB GGB TGB CTT TTG BGG GGG BCB CBG BTG TCT GGG CBT TGC CBG GTC CTG GGB BCB GBG CCC CGB GCB GGB GCB GGG GCG GCG GCC GGG GCC TGC TGG GBG CCB TBG CGB GGC TGB G-3' 10 (FRAG. NO:1073) (SEQ. ID NO:1083)

Human Vascular Cell Adhesion Molecule 1 (VCAM-1) Anti-sense Oligonucleotide

- CCG CIT TIT TCT TC GTC TTT GTT GTT TTC TCT TCC TTG CTG BGC BBG BTB TCT BGB TTC TGG GGT GGT CTC GBT 15 TTT BBBB GCT TGB GBB GCT GCB BBC BTT BTC CBB BGT BTB TTT GBG GCT CCB BGG BTC BCG BCC BTC TTC CCB GGC BTT TTB BGT TGC TGT CGT-3' (FRAG. NO: 1735) (SEQ. ID NO: 1746) 5'-C TGT CGT-3' (FRAG. NO:1736) (SEQ. ID NO:1747) 5'-TGC TTC TTC C-3' (FRAG. NO:1737) (SEQ. ID NO:1748)
- HSVCAM1AS1: 5'-CCT CTT TTC TGT TTT TCC C-3' (FRAG. NO:1074) (SEQ. ID NO:1084) 20 HSVCAM1AS2: 5'-CTC TGC CTT TGT TTG GGT TCG-3' (FRAG, NO:1075) (SEQ. ID NO:1085) HSVCAMIAS3: 5'-CTT CCT TTC TGC TTC TTC C-3' (FRAG. NO:1076) (SEQ. ID NO:1086) HSVCAM1AS4: 5'-CTG TGT CTC CTG TCT CCG CTT TTT TCT TC-3' (FRAG. NO:1077) (SEQ. ID NO:1087) HSVCAM1ASS: 5'-GTC TTT GTT GTT TTC TCT TCC TTG-3' (FRAG. NO:1078) (SEQ. ID NO:1088)
- CTG BGC BBG BTB TCT BGB TTC TGG GGT GGT CTC GBT TTT BBBB GCT TGB GBB GCT GCB BBC BTT BTC CBB BGT BTB TTT GBG GCT CCB BGG BTC BCG BCC BTC TTC CCB GGC BTT TTB BGT TGC TGT CGT (FRAG. NO:1079) (SEQ. ID NO:1089) 25

Human Endothelial Leukocyte Adhesion Molecule (ELAM-1) Anti-sense Oligonucleotide Fragments

- 5'-BBG TGB GBG CTG BGB GBB BCT GTG BBG CBB TCB TGB CTT CBB GBG TTC TTT TCB CCC GTT CTT GGC TTC TTC TGT CTG G GTT CTT GGC TTC TTC TGT CCG T TGG CTT CTC GTT GTC CC TGT GGG CTT CTC GTT GTC CC CCC TTC GGG 30 GGC TGG TGG GGC CGT CCT TGC CTG CTG G (FRAG. NO: 1738) (SEQ. ID NO: 1749) 5'-CCT TGC CTG CTG G-3' (FRAG. NO: 1739) (SEQ. ID NO: 1750)
 - 5'-GTT GTC CC-3' (FRAG. NO: 1740) (SEQ. ID NO:1751) 5'-GTT CTT GGC TTC TTC TGT C-3' (FRAG. NO:1080) (SEQ. ID NO:1090)
- 35 5'-GGC TGG TGG-3' (FRAG. NO:1083) (SEQ. ID NO:1093) 5'-CGT TGG CTT CTC GTT GTC CC-3' (FRAG. NO:1081) (SEQ. ID NO:1091) 5'-TGT GGG CTT CTC GTT GTC CC-3' (FRAG. NO:1082) (SEQ. ID NO:1092) 5'-CCC TTC GGG GGC TGG TGG-3' (FRAG. NO:1083) (SEQ. ID NO:1093)
- 5'-GGC CGT CCT TGC CTG CTG G-3' (FRAG. NO:1084) (SEQ. ID NO:1094) 40

Human P Selectin Fragments

- CTT TTC-3' (FRAG. NO: 1741) (SEQ. ID NO: 1752) 5'-TCC TTT CTT TTC-3' (FRAG. NO: 1742) (SEQ. ID NO: 1753)
- 5'-CTC CTT TT-3' (FRAG. NO:1743) (SEQ. ID NO:1754) 45
 - 5'-TTG CTG TTT TTT CTC CTT CTT CTC TCC TTT CTT TTC-3' (FRAG. NO:1086) (SEQ. ID NO:1096)

Human Endothelial Monocyte Activating Factor Anti-sense Oligonucleotide Fragments

CTT TTC-3' (FRAG. NO: 1744) (SEQ. ID NO: 1755) 50 5' -CC TTT CTT TTC (FRAG. NO: 1745) (SEQ. ID NO: 1756) 5'-CTG TTC CTC CTT TT-3' (FRAG. NO:1746) (SEQ. ID NO:1757) 5'-TIG CTG TTT TTT CTC CTT CTT CTC TCC TTT CTT TTC-3' (FRAG. NO:1088) (SEQ. ID NO:1098)

Human IL3* Anti-sense Oligonucleotide Fragments

GGB GCB GGB GGB GGG GGC GGC TCB TGT TTG GBT CGG CBG GBG GCB CTC (FRAG. NO: 1747) (SEQ. ID NO:

5'-G GBG GCB CTC-3' (FRAG. NO: 1748) (SEQ. ID NO: 1759) 5'-GT GGG GCT CTG-3' (FRAG. NO:1749) (SEQ. ID NO:1760) HUMIL3AAS1: 5'-CTC TGT CTT GTT CTG GTC CTT CGT GGG GCT CTG-3' (FRAG. NO:1089) (SEQ. ID NO:1099) HUMIL3AAS2: 5'-TGT CGC GTG G GTG CGG CCG TGG CC-3' (FRAG. NO:1090) (SEQ. ID NO:1100) NO:1091) (SEQ. ID NO:1101) 10

Human IL3 Receptor Anti-sense Oligonucleotide Fragments

GCG CTT GTC GTT TTG GGG CCG GCT TTG CCC GCC TCC CGG CGC CTG GCC CGG CC TTC CTG GGC TGC GTG CGC GTT CTG TTC TTC TTC CTG GCT CTG GGG TGT CCT GGC CTT CGT GGT TCC TCT TCC TTC GTT TGC CGT CCG CGG GGG CCC 15 CCT CCT CCT GGT CGC GCT TGT CGT TTT GG GGC CGG CTT TGC CCG CCT CCC GGC GCC TGG CCC GGC CTT CCT GGG CTG CGT GCG CGT TCT CTT CTT CCT GGC GCA GGA GAC AGG GCA GGG CGA TCA GGA GCA GCG TGA GCC AAA GGA GGA CCA TCG GGA ACG CAG CTC CGG AAC GCA GGA CAG AGG TGC C GC BGG BGB CBG GGC BGG GCG BTC BGG

BGC BGC GTG BGC CBB BGG BCC BTC GGG BBC GCB GCT CCG GBB CGC BGG BCB GBG GTG CC-3' (FRAG. NO: 1750) 20 (SEO. ID NO: 1761)

GBG GTG CC-3' (FRAG. NO: 1751) (SEQ. ID NO: 1762)

5'- GCC CCG C-3' (FRAG. NO:1752) (SEQ. ID NO:1763)

5'-TCTGGGGTGTCCTG (FRAG. NO:1092) (SEQ. ID NO:1102)

5'-GCCTTCGTGGTTCC (FRAG. NO:1093) (SEQ. ID NO:1103) 25

5'-TCTTCCTTCGTTTGC (FRAG. NO:1094) (SEQ. ID NO:1104)

5'-CGTCCGCGGGGCCCCCGGGCCT (FRAG. NO:1095) (SEQ. ID NO:1105)

5'-GGC TGC GCT CCT GCC CCG C (FRAG. NO:1096) (SEQ. ID NO:1106)

5'-CTCTTTCCCGGGCTCTT (FRAG. NO:1097) (SEQ. ID NO:1107)

5'-GCGCTGGGGGGTGCTCC (FRAG. NO:1098) (SEQ. ID NO:1108) 30

5'-CGTGTGTTTGCGCCCTCCTGGTCGC (FRAG. NO:1099) (SEQ. ID NO:1109)

5'-GCTTGTCGTTTTGG (FRAG. NO:1100) (SEQ. ID NO:1110)

5'-GGCCGGCTTTGCCCGCCTCCC (FRAG. NO:1101) (SEQ. ID NO:1111)

5'-GGCGCCTGGCCCGGCC (FRAG. NO:1102) (SEQ. ID NO:1112)

5'-TTCCTGGGCTGCGTGCGC (FRAG. NO:1103) (SEQ. ID NO:1113) 35

5'-GTTCTGTTCTTCCTGGC (FRAG. NO:1104) (SEQ. ID NO:1114)

GCB GGB 5'-CBG BGG TGC C (FRAG. NO:1105) (SEQ. ID NO:1115)

Human IL-4 Anti-sense Oligonucleotide Fragments

5'-CTC TGG TTG GCT TCC TTC GCC GGC BCB TGC TBG CBG GBB GBB CBG BGG GGG BBG CBG TTG GGB GGT GBG BCC 40 CBT TBB TBG GTG TCG B-3' (FRAG. NO: 1753) (SEQ. ID NO: 1764) 5'-GCC GGC BCB-3' (FRAG. NO: 1754) (SEQ. ID NO: 1765) 5'-T TCC TTC-3' (FRAG. NO:1755) (SEQ. ID NO:1766)

5'-CTC TGG TTG GCT TCC TTC-3' (FRAG. NO:1106) (SEQ. ID NO:1116) 5'-GCCGGCBCBTGCTBGCBGGBGGBGGGGGGBGGCBGTTGGGBGCCCBTTBBTBGGTGTCGB-3' (FRAG. NO:1107) 45 (SEO. ID NO:1117)

Human IL4 Receptor Anti-sense Oligonucleotide Fragment

5'-TCT GCC CTG TCC GCC GGC TCT TCG GTG GCT CGG CCC CGC TCC TTG TCT TGC CGC GGG TTG GTT CCT GGG CCT 50 CCC CTC GGG CGG CTG CGG GCG CTC GTG CCT GGT CCG CTC CCT GGG GGT GCT CCT TCC CTT TCC CCG CTC GTG TGG GCT TCC CTG TGC CCC TTT CCT CTG GGT CCC CCT CCC GTT CCA AGC TGC ACC GCA CAG ACC GGC GCT ACA GGA CAG AGC CAG GCA AGC ACC CAT GGG GAT CCA GGC CCA GCT GTT CCB BGC TGC BCC GCB CBG BCC GGC GCT BCB GGB CBG CBG GCB BGC BCC CBT GGG GBT CCB GGC CCB GCT G -3' (FRAG. NO: 1756) (SEQ ID NO: 1767) 55

5'-TCTGCGC-3' (FRAG. NO: 1757) (SEQ ID NO: 1768)

5'-CCT GCT CCT GGG G (FRAG. NO:1758) (SEQ. ID NO:1769) 5'-TCTGCGCGCCCCTGCTCC (FRAG. NO:1108) (SEQ. ID NO:1118)

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5'-CGCCCGGCTTCTCT (FRAG. NO:1109) (SEQ. ID NO:1119)
      5'-CGTGTGGGCTTCGG (FRAG. NO:1110) (SEQ. ID NO:1120)
      5'-CCCCGCGCCTCCGTTGTTCTC (FRAG. NO:1111) (SEQ. ID NO:1121)
      5'-TGCTCGCTGGGCTTG (FRAG. NO:1112) (SEQ. ID NO:1122)
      5'-GGTTTCCTGGGGCCCTGGGTTTC (FRAG. NO:1113) (SEQ. ID NO:1123)
5
       5'-TCTGCCGGGTCGTTTTC (FRAG. NO:1114) (SEQ. ID NO:1124)
       5'-GGGTGCTGCG (FRAG. NO:1115) (SEQ. ID NO:1125)
       5'-CTTGGTGCTGGGGCTCC (FRAG. NO:1116) (SEQ. ID NO:1126)
       5'-GGCGGCTGCGGGCTGGGTTGGG (FRAG. NO:1117) (SEQ. ID NO:1127)
       5'-CTTGGCTGGTTCCTGGCCTCGGG (FRAG. NO:1118) (SEQ. ID NO:1128)
10
       5'-CCTCCTCCTCCTCCTCCTCCTCTTTTTCTTCCTCT (FRAG. NO:1119) (SEQ. ID NO:1129)
       5'-TCCCTGCTGCTCTC (FRAG. NO:1120) (SEQ. ID NO:1130)
       5'-TGCCCTCCCTTCCCTCCTGG (FRAG. NO:1121) (SEQ. ID NO:1131)
       5'-GGTGCCTCCTTGGGCCCTGC (FRAG. NO:1122) (SEQ. ID NO:1132)
       5'-GGCTGCTCCTTGCCCC (FRAG. NO:1123) (SEQ. ID NO:1133)
15
       5'-CTCTGGGTCGGGCTGGC (FRAG. NO:1124) (SEQ. ID NO:1134)
       5'-GGGGCGTCTCTGTGC (FRAG. NO:1125) (SEQ. ID NO:1135)
       5'-CTGGCCTGGGTGCC (FRAG. NO:1126) (SEQ. ID NO:1136)
       5'-GCCTCTCCTGGGGGGGGGGCTCCCTGTCC (FRAG. NO:1127) (SEQ. ID NO:1137)
       5'-CCTTTTCCCCCGGCTCC (FRAG. NO:1128) (SEQ. ID NO:1138)
20
       5'-GTGGGGGCTTTGGC (FRAG. NO:1129) (SEQ. ID NO:1139)
       5'-GGG GGT CTG TGG CCT GCT CCT GGG G (FRAG. NO:1130) (SEQ. ID NO:1140)
       5'-AGGGGTCTGGGGCCCTC (FRAG. NO:1131) (SEQ. ID NO:1141)
       5'-TITTGGGGGTCTGGCTTG (FRAG. NO:1132) (SEQ. ID NO:1142)
       5'-GCCTGGCTGCCTTCC (FRAG. NO:1133) (SEQ. ID NO:1143)
25
       5'-GGGGCCTGCCGTGGGGC (FRAG. NO:1134) (SEQ. ID NO:1144)
       5'-TGTCCTCTGTTGCTCCCCTT (FRAG. NO:1135) (SEQ. ID NO:1145)
       5'-TGCCTGCTGTCTGG (FRAG. NO:1136) (SEQ. ID NO:1146)
        5'-GGTTCCCGCCTTCCCT (FRAG. NO:1137) (SEQ. ID NO:1147)
        5'-GTT CCC AGA GCT TGC CAC CTG CAG GAC CAG GCA GCT CAC AGG GAA CAG GAG CCC AGA GCA AAG CCA CCC
30
        CAT TGG GAG ATG CCA AGG CAC CAG GCT G (FRAG. NO:1138) (SEQ. ID NO:1148)
        5'-GTT CCC BGB GCT TGC CBC CTG CBG CBG GBC CBG GCB GCT CBC BGG GBB CBG GCB BBG CCB CCC
        CBT TGG GBG BTG CCB BGG CBC CBG GCT G-3' (FRAG. NO:1139) (SEQ. ID NO:1149)
        Human IL5* Anti-sense Oligonucleotide Fragments
        5'-TCCCTGTTTC CCCCCTTTCG TTCTGCGTTT GCCTTTGGCG TTTTTTGTTT GTTTTCTCTC TCCGTCTTTC TTCTCCCCT
 35
        GTGGGBBTTT CTGTGGGGBT GGCBTBCBCG TBGGCBGCTC CBBGBGCTBG CBBBCTCBBB TGCBGBBGCB TCCTCBTGGC
        TCTGBBBCGG TGGGAATTTC TGTGGGGBTG GCATACACGT AGGCAGCTCC AAGAGCTAGC AAACTCAAAT
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- GCAGAAGCATC CTCATGGCTC TGAAACG-3' (FRAG. NO: 1759) (SEQ. ID NO: 1770)
 5'-GCC CCG GG-3' (FRAG. NO: 1760) (SEQ. ID NO: 1771)
 5'-G GGT TTC T-3' (FRAG. NO: 1761) (SEQ. ID NO: 1772)
- 40
 - 5'-GTG GGG BTG GC-3' (FRAG. NO: 1762) (SEQ. ID NO:1773_)
 5'-CCB BGB GCT BGC-3' (FRAG. NO: 1763) (SEQ. ID NO: 1774)
 - 5'-TCC CTG TTT CCC CCC TTT-3' (FRAG. NO:1140) (SEQ. ID NO:1150)
 - 5'-CGT TCT GCG TTT GCC TTT GGC-3' (FRAG. NO:1141)(SEQ. ID NO:1151)
- 5'-GTT TTT TGT TTG TTT TCT-3' (FRAG. NO:1142)(SEQ. ID NO:1152) 45
- 5'-CTC TCC GTC TTT CTT CTC C-3' (FRAG. NO:1143) (SEQ. ID NO:1153)
 - 5'-CCT CCT GCC TGT GTC CCT GCT CCC C-3' (FRAG. NO:1144) (SEQ. ID NO:1154)
 - 5'-GAG GGT TTC TGG CTT CCT CTC T-3' (FRAG. NO:1145) (SEQ. ID NO:1155)
 - 5'-TGT CTC TCT GTC CTT TTG TT-3' (FRAG. NO:1146) (SEQ. ID NO:1156)
- 5'-TGT TGT GCG GCC TGG TGC TGC CCT GCC CCG GG-3' (FRAG. NO:1147) (SEQ. ID NO:1157) 50 5'-GTG GGA ATT TCT GTG GGG BTG GCA TAC ACG TAG GCA GCT CCA AGA GCT AGC AAA CTC AAA TGC AGA AGC ATC CTC ATG GCT CTG AAA CG-3' (FRAG. NO: 1764) (SEQ. ID NO: 1775) 5'-GTG GGB BTT TCT GTG GGG BTG GCB TBC BCG TBG GCB GCT CCB BGB GCT BGC BBB CTC BBB TGC BGB BGC BTC CTC BTG GCT CTG BBB CG-3' (FRAG. NO:1148) (SEQ. ID NO:1158)

Human IL-5 Receptor Anti-sense Oligonucleotide Fragments 55

5'-CTCAGTGGCC CCCAAAAGGA TGAGTAATAC ATGCGCCACG ATGATCATAT CCTTTTTACT ATGAGGCCGT GCTGTGCCCT GCCTCTCTGC-3' (FRAG. NO: 1765) (SEQ. ID NO: 1776)

5'-CCG TGT C-3' (FRAG. NO: 1766) (SEQ. ID NO: 1777) 60

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5'-GCCCTGCC-3' (FRAG. NO: 1767) (SEQ. ID NO: 1778)
      5'-CCG TGT CTG TCG TGT CT-3' (FRAG. NO:1149) (SEQ. ID NO:1159)
      5'-TTCCTTTGCTCTTG-3' (FRAG. NO:1150) (SEQ. ID NO:1160)
      5'-GTGTGTCTTTGCTGT-3' (FRAG. NO:1151) (SEQ. ID NO:1161)
5'-GCCCTGCCTCTCTGC-3' (FRAG. NO:1152) (SEQ. ID NO:1162)
5
      5'-CT CBGTGGCCCC CBBBBGGBTG BGTBBTBCBT GCGCCBCGBT GBTCBTBTCC TTTTTBCTBT GBGG (FRAG. NO: 1768)
       (SEQ. ID NO: 1779)
      Human IL-6 Receptor Fragments
      5'-GGGGGTGGCT TCCTGCCGCG TCTCTGGGCC GTCCCGTCCC TCGGCCCCGC GCCGCGCTCG GCTCCTCTCC CTCTGGCCCG
       10
       TGCTGGCTGC CCTGCTGGCC GCGCCGGGGC CTGTCCGCCT CTGCGGGCGC TGTCTCCTGG CTTGTCTTCC GGCTCTTCTG
       GGGGGTTTCT GGCCGTGGGG GTCTTGCCTG GCCTCCGGGC TCCTGCTTGT CTTGCCTTCC TTCTCTGGTC GGTTGTGGCT
       CTGCTTCTCG CTCTCCTTTG TGGGGCCCTC CCTGCTGCTC TTGGTTTTGG GCTTTTTTTC TCTTCCTCCT TTTTCGTGCG
15
       TGGGCCTCCG CACGCCTCTT GCCACCTCCT GCGCAGGGCA GCGCCTTGG GGCCAGCGCC GCTCCCGGCG CGGCCAGCAG
       GGCAGCCAGC AGCGCGCAGC CGACGGCCAG CATGCTTCCT CCTCGGCTAC CACTCCATGG TCCCGCAGAG GCGGACAGGC
       GCBCGCCTC TTGCCBCCTC CTGCGCBGGG CBGCGCCTTG GGGCCBGCGC CGCTCCCGGC GCGGCCBGCB GGGCBGCCBG
       CBGCGCGCBG CCGBCGGCCB GCBTGCTTCC TCCTCGGCTB CCBCTCCBTG GTCCCGCBGB GGCGGBCBGG C-3'
        (FRAG. NO: 1769) (SEQ. ID NO: 1780)
20
       5'-CCCGGCGC-3' (FRAG. NO:1184) (SEQ. ID NO:1194)
       5'-GGCCBGCBGG-3' (FRAG. NO:1186) (SEQ. ID NO:1196)
       5'-GCBGCCBGCBGCG-3' (FRAG. NO: 1770) (SEQ. ID NO: 1781)
       5'-C GCBGCCGBCGGCC -3' (FRAG. NO: 1771) (SEQ. ID NO: 1782)
       5'-GGGGGTGGCTTCCTGCC3'- (FRAG. NO:1153) (SEQ. ID NO:1163)
5'-GCGTCTCTGGGCCGTCCC-3' (FRAG. NO:1154) (SEQ. ID NO:1164)
25
        5'-GTCCCTCGGCCCCGCGCCGCGCTCGGCTCCTCTCCC-3' (FRAG. NO:1155) (SEQ. ID NO:1165)
        5'-TCTGGCCCGGCTC-3' (FRAG. NO:1156) (SEQ. ID NO:1166)
        5'-GGCGCTGCCCTGCGC-3' (FRAG. NO:1158) (SEQ. ID NO:1168)
30
        5'-GCGGCGCTGGCCCC-3' (FRAG. NO:1159) (SEQ. ID NO:1169)
        5'-TGCTGGCCGTCGGCTGCTGCTGCCCT-3' (FRAG. NO:1160) (SEQ. ID NO:1170)
        5'-GCTGGCCGCCGGG-3' (FRAG. NO:1161) (SEQ. ID NO:1171)
        5'-GCCTGTCCGCCTCTGCGGG-3' (FRAG. NO:1162) (SEQ. ID NO:1172)
        5'-CGCTGTCTCCTGGC-3' (FRAG. NO:1163) (SEQ. ID NO:1173)
5'-TTGTCTTCCGGCTCT-3' (FRAG. NO:1164) (SEQ. ID NO:1174)
 35
        5'-TCTGCTGGGGTGGG-3' (FRAG. NO:1165) (SEQ. ID NO:1175)
        5'-GCTGGGCCGGCCCGGT-3' (FRAG. NO:1166) (SEQ. ID NO:1176)
5'-GCTGGGGCTCCTCGGGGGG-3' (FRAG. NO:1167) (SEQ. ID NO:1177)
        5'-GGGGGCTCTTCCGG-3' (FRAG. NO:1168) (SEQ. ID NO:1178)
5'-GCTGTCTCCCTCCGGG-3' (FRAG. NO:1169) (SEQ. ID NO:1179)
5'-GCGGGGGTTTCTGGCC-3' (FRAG. NO:1170) (SEQ. ID NO:1180)
 40
        5'-GTGGGGGTCTTGCC-3' (FRAG. NO:1171) (SEQ. ID NO:1181)
        5'-TGGCCTCCGGGCTCC-3' (FRAG. NO:1172) (SEQ. ID NO:1182)
        5'-TGCTTGTCTTGCCTTCC-3' (FRAG. NO:1173) (SEQ. ID NO:1183)
5'-TCTGGTCGGTTGTGGCTCG-3' (FRAG. NO:1174) (SEQ. ID NO:1184)
 45
        5'-GGGCTCCGTGGGTCCCTGGC-3' (FRAG. NO:1175) (SEQ. ID NO:1185)
5'-GCCCGTTTGTGTTTTGTC-3' (FRAG. NO:1176) (SEQ. ID NO:1186)
         5'-TTTTCCCCTGGCGT-3' (FRAG. NO:1177) (SEQ. ID NO:1187)
         5'-CCCTGTGCCCCTCTCCTCCTCCTCTCTCTC-3' (FRAG. NO:1178) (SEQ. ID NO:1188)
  50
         5'-GCTCTCCTTTGTGGG-3' (FRAG. NO:1179) (SEQ. ID NO:1189)
         5'-GCCCTCCCTGCTGCT-3' (FRAG. NO:1180) (SEQ. ID NO:1190)
         5'-CTTGGTTTTGGGCT-3' (FRAG. NO:1181) (SEQ. ID NO:1191)
         5'-TTTTTTCTCTTCCTCCTTTTTC-3' (FRAG. NO:1182) (SEQ. ID NO:1192)
         5'-GTGCGTGGGCCTCC-3' (FRAG. NO:1183) (SEQ. ID NO:1193)
  55
         CGCAGCCGACGGCCAGCATGCTTCCTCCTCGGCTACCACTCCATGGTCCCGCAGAGGCGGACAGGC-3'
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GCBGCCGBCGGCCBGCBTGCTTCCTCCTCGGCTBCCBCTCCBTGGTCCCGCBGBGGCCGBCBGGC-3'

(FRAG. NO:1185) (SEQ. ID NO:1195)

60

(FRAG. NO:1187) (SEQ. ID NO:1197)

Human IL-6 Anti-sense Oligonucleotide Fragments

5'-GGGGGTGGCT TCCTGCCGCG TCTCTGGGCC GTCCCGTCCC TCGGCCCCGC GCCGCGCTCG GCTCCTCTCC CTCTGGCCCG TGCTGGCTGC CCTGCTGGCC GCGCCGGGGC CTGTCCGCCT CTGCGGGCGC TGTCTCCTGG CTTGTCTTCC GGCTCTTCTG 5 GGGGGTTTCT GGCCGTGGGG GTCTTGCCTG GCCTCCGGGC TCCTGCTTGT CTTGCCTTCC TTCTCTGGTC GGTTGTGGCT CTGCTTCTCG CTCTCTTTG TGGGGCCCTC CCTGCTGCTC TTGGTTTTGG GCTTTTTTTC TCTTCCTCCT TTTTCGTGCG TGGGCCTCC GCACGCCTCT TGCCACCTCC TGCGCAGGGC AGCGCCTTGG GGCCAGCGCC GCTCCCGGCG CGGCCAGCAG

10 GGCAGCCAGC AGCGCGCAGC CGACGGCCAG CATGCTTCCT CCTCGGCTAC CACTCCATGG TCCCGCAGAG GCGGACAGGC GCBCGCCTC TTGCCBCCTC CTGCGCBGGG CBGCGCCTTG GGGCCBGCGC CGCTCCCGGC GCGGCCBGCB GGGCBGCCBG CBGCGCGCBG CCGBCGGCCB GCBTGCTTCC TCCTCGGCTB CCBCTCCBTG GTCCCGCBGB GGCGGBCBGG C-3' (FRAG. NO:1772) (SEQ. ID NO:1783)

5'-GGGGCBGG-3' (FRAG. NO:1773) (SEQ. ID NO:1784) 15

- 5'-GBBGGCBG CBGGC-3' (FRAG. NO:1774) (SEQ. ID NO:1785) 5'-CCBGGBGCBG CCCC-3' (FRAG. NO:1775) (SEQ. ID NO:1786)
- 5'-BGGG BGBBGGCBBC-3' (FRAG. NO:1776) (SEQ. ID NO:1787)
- 5'-GCT TCT CTT TCG TTC CCG GTG GGC TCG-3' (FRAG. NO:1188) (SEQ. ID NO:1198)
- 5'-GTG GCT GTC TGT GTG GGG CGG CT-3' (FRAG. NO:1189) (SEQ. ID NO:1199) 20

 - 5'-GTG CCT CTT TGC TGC TTT C-3' (FRAG. NO:1190) (SEQ. ID NO:1200)
 5'-GAT TCT TTG CCT TTT TCT GC-3' (FRAG. NO:1191) (SEQ. ID NO:1201)
 - 5'-CTCCTGGGGG TBCTGGGGCB GGGBBGGCBG CBGGCBBCBC CBGGBGCBGC CCCBGGGBGB BGGCBBCTGG BCCGBBGGCG CTTGTGGBGB BGGBGTTCBT BGCTGGGCTC CTGGBGGGGB GBTBGBGC-3' (FRAG. NO:1777) (SEQ. ID NO:1788)

Human Monocyte-derived Neutrophil Chemotactic 25 Factor Anti-sense Oligonucleotide Fragments

5'-GGGGTGGBBB GGTTTGGBGT BTGTCTTTBT GCBCTGBCBT CTBBGTTCTT TBGCBCTCCT TGGCBBBBCT GCBCCTTCBC BCBGBGCTGC BGBBBTCBGG BBGGCTGCCB BGBGBGCCBC GGCCBGCTTG GBBGTCBTGT TTBCBCBCBG TGBGBTGGTT CCTTCCGGGC TTGTGTGCTC TGGTTCCTTC CGGTGGTTTC TTCCTGGCTC TTGTCCTTTC TCTTGG CCCT

TGGC-3' (FRAG. NO:1778) (SEQ. ID NO: 1789) 30

- 5'-GGBGT BTG-3' (FRAG. NO:1779) (SEQ. ID NO: 1790)
- 5'-GCBCTGBCBT CT-3' (FRAG. NO:1780) (SEQ. ID NO:1791)
- 5'-CCG GTG G-3' (FRAG. NO:1781) (SEQ. ID NO: 1792)
- 5'-GG CCC TTG GC-3' (FRAG. NO:1782) (SEQ. ID NO: 1793)
- 35
- 5'-GCT TGT GTG CTC TGC TGT CTC T-3' (FRAG. NO:1192) (SEQ. ID NO:1202) 5'-TGG TTC CTT CCG GTG GTT TCT TCC TGG CTC TTG TCC T-3' (FRAG. NO:1193) (SEQ. ID NO:1203)
 - 5'-TTC TCT TGG CCC TTG GC-3' (FRAG. NO:1194) (SEQ. ID NO:1204)
 - 5'-GGGGTGGBBB GGTTTGGBGT BTGTCTTTBT GCBCTGBCBT CTBBGTTCTT TBGCBCTCCT TGGCBBBBCT GCBCCTTCBC BCBGBGC-3' (FRAG. NO:1783) (SEQ. ID NO: 1794)

Human Neutrophil Elastase (Medullasin) Anti-sense Oligonucleotide Fragments 40

5'-GGGCTCCCGC CGCGBGBGGT TBTGGGCTCC CBGGBCCBCC CGCBCCGCGC GGBCGTTTBC BTTCGCCBCG CBGTGCGCGG CCGBCBTGBC GBBGTTGGCC GCBBTCBGGG TGGCGCCGCB GBBGTGGCCT CCGCGCBGCT GCBGGGBCBC CBTGBBGGGC CBCGCGTGGG GCCGCGCTCG CCGGCCCCC BCBBTCTCCG BGGCCBGCGC GGTGCCCCCC BGCBGCBBGG CCGGCBGGBC BCBGGCGBGG BGBCBCGCGB GTCGGCGGCC GBGGGTCBTG GTGGGGCTGG GGCTCCGGGG TCTCTGCCCC TCCGTGCTGG

- TGGGGCTGGG GCTCCGGGG TCTCTGCCCC TCCGTGCCGC GTGGGGGCCGC GCTCGCCGGC CCCCCCTGC CGGGTGGGCT 45 CCCGCCGCG GCCGCCTGC CGGCCCCTCG TGGCTCCTGC TGGCCGGGTCCCGG GGGTGGGGGCG CGBGTCGGCG GCCGBGGGTC-3' (FRAG. NO:1784) (SEQ. ID NO: 1795) 5'-GG TGG GGC-3' (FRAG. NO:1785) (SEQ. ID NO: 1796)

 - 5'-G GGG CCG -3' (FRAG. NO:1786) (SEQ. ID NO:1797)
- 5'- GGC CGG GTC CGG G-3' (FRAG. NO:1787) (SEQ. ID NO: 1798) 50
- 5'-TGG TGG GGC TGG GGC TCC GGG GTC TCT GCC CCT CCG TGC-3' (FRAG. NO:1195) (SEQ. ID NO:1205)
 - 5'-CGC GTG GGG CCG CGC TCG CCG GCC CCC C-3' (FRAG. NO:1196) (SEQ. ID NO:1206)
 - 5'-CCT GCC GGG TGG GCT CCC GCC GCG-3' (FRAG. NO:1197) (SEQ. ID NO:1207)
 - 5'-CGC CGG CCT GCC GGC CCC TC-3' (FRAG. NO:1198) (SEQ. ID NO:1208)
- 5'-GTG GGT CCT GCT GGC CGG GTC CGG GTC CCG GGG GTG GGG-3' (FRAG. NO:1199) (SEQ. ID NO:1209) 55
 - 5'-CGC GBG TCG GCG GCC GBG GGT C-3' (FRAG. NO:1200) (SEQ. ID NO:1210)
 - 5'-GGCTCCCGC CGCGBGBGGT TBTGGGCTCC CBGGBCCBCC CGCBCCGCGC GGBCGTTTBC BTTCGCCBCG CBGTGCGCGG

CCGBCBTGBC GBBGTTGGCC GCBBTCBGGG TGGCGCCGCB GBBGTGGCCT CCGCGCBGCT GCBGGGBCBC CBTGBBGGGC CBCGCGTGGG GCCGCCTCG CCGGCCCCCC BCBBTCTCCG BGGCCBGCGC GGTGCCCCCC BGCBGCBBGG CCGGCBGGBC BCBGGCGBGG BGBCBCGCGB GTCGGCGGCC GBGGGTCBTG GTGGGGCTGG GGCTCCGGGG TCTCTGCCCC TCCGTGC-3' (FRAG. NO:1788) (SEQ. ID NO: 1799)

Human Neutrophil Oxidase Factor Anti-sense Oligonucleotide Fragments

5'-CGGGBGTGGG GGTCCTGGBC GGCBCTGBBG GCBTCCBGGG CTCCCTTCCB GTCCTTCTTG TCCGCTGCCB GCBCCCCTTC BTTCCBGBGG CTGBTGGCCT CCBCCBGGGB CBTGBTTBGG TBGBBBCTBG GBGGCCGGCC TCCBCCBGGG BCBTGGTCCT CTGGCCTGGT GCTCTCTCGT GCCCTTTCCC TTGGGTGTCT TGTTTTTGTG GCCTCCBCCB GGGBCBTG-3' (FRAG. NO:1789)

- 10 (SEQ. ID NO: 1800)
 - 5'-CGGGBGTGGG GG-3 ' (FRAG.NO:1790) (SEQ. ID NO: 1801)
 - 5'-GCCBGCBCCCC-3' (FRAG.NO:1791) (SEQ. ID NO: 1802)
 - 5'-C CBC CBG-3' (FRAG.NO:1792) (SEQ. ID NO: 1803)
 - 5'-GGC CTC CBC CBG GGB CBT G-3' (FRAG. NO:1201) (SEQ. ID NO:1211)
- 5'-GTC CTT CTT GTC CGC TGC C -3' (FRAG. NO:1202) (SEQ. ID NO:1212) 15
 - 5'-TCT CTG GGG TTT TCG GTC TGG GTG G-3 (FRAG. NO:1203) (SEQ. ID NO:1213) 5'-GCT TTC CTC CTG GGG CTG CTG CTG-3' (FRAG. NO:1204) (SEQ. ID NO:1214)
 - 5'-GGC TCT TCT TTT TGT TTC TGG CCT GGT G-3' (FRAG. NO:1205) (SEQ. ID NO:1215)
 - 5'-CTC TCT CGT GCC CTT TCC-3' (FRAG. NO:1206) (SEQ. ID NO:1216)
- 20
- 5'-CTT GGG TGT CTT GTT TTT GT-3' (FRAG. NO:1207) (SEQ. ID NO:1217) 5'-GGC CTC CBC CBG GGB CBT G-3' (FRAG. NO:1208) (SEQ. ID NO:1218)
 - 5'-CGGGBGTGGG GGTCCTGGBC GGCBCTGBBG GCBTCCBGGG CTCCCTTCCB GTCCTTCTTG TCCGCTGCCB GCBCCCCTTC BTTCCBGBGG CTGBTGGCCT CCBCCBGGGB CBTGBTTBGG TBGBBBCTBG GBGGCC-3' (FRAG. NO:1793) (SEQ. ID NO:

Human Cathepsin G Anti-sense Oligonucleotide Fragments 25

- 5'-CCCTCCBCBT CTGCTCTGBC CTGCTGGBCT CTGGBTCTGB BGBTBCGCCB TGTBGGGGCG GGBGTGGGGC CTGCTCTCCC GGCCTCCGBT GBTCTCCCCT GCCTCBGCCC CBGTGGGTBG GBGBBBGGCC BGCBGBBGCB GGBGTGGCTG CBTCTTTCCT GGTGGGGCCT GCTCTCCCGG CCTCCGTGTG TTGCTGGGTG TTTTCCCGTC TCTGGTCTGC CTTCGGGGGT CGT-3' (FRAG. NO:1794) (SEQ. ID NO: 1805)
- 5'-GBBGBTBCGCC-3' (FRAG. NO:1795) (SEQ. ID NO: 1806) 30
 - 5'-CBGCCCCBG-3' (FRAG. NO:1796) (SEQ. ID NO: 1807)
 - 5'-TCC CGT CTC TGG-3' (FRAG. NO:1797) (SEQ. ID NO: 1808)
 - 5'-GTG GGG CCT CCC CCG GCC TCC G-3' (FRAG. NO:1209) (SEQ. ID NO:1219)
 - 5'-TGT GTT GCT GG GTG TTT TCC CGT CTC TGG-3' (FRAG. NO:1210) (SEQ. ID NO:1220)
- 5'-TCT GCC TTC GGG GGT CGT-3' (FRAG. NO:1211) (SEQ. ID NO:1221) 35
- 5'-CCCTCCBCBT CTGCTCTGBC CTGCTGGBCT CTGGBTCTGB BGBTBCGCCB TGTBGGGGCG GGBGTGGGGC CTGCTCTCCC GGCCTCCGBT GBTCTCCCCT GCCTCBGCCC CBGTGGGTBG GBGBBBGCC BGCBGBBGCB GGBGTGGCTG-3 NO:1798) (SEQ. ID NO: 1809)

Human Defensin 1 Anti-sense Oligonucleotide Fragments

- 5'-CCGGGGCTGC BGCBBCCTCB TCBGCTCTTG CCTGGBGTGG CTCBGCCTGG GCCTGCBGGG CCBCCBGGBG BBTGGCBGCB BGGBTGGCGB GGGTCCTCBT GGCTGGGGTC BCBGBTCCTC TBGCTBGGCB GGGTGBCCBG BGBGGGC GGG TCC TCB TGG 40 CTG GGG GCC TGG GCC GCT CTT GCC TGG BGT GGC TC GCC CBG BGT CTT CCC TGG T-3' (FRAG.NO:1799) (SEQ. ID NO: 1810)
 - 5'-CCGGGGC-3' (FRAG.NO:1800) (SEQ. ID NO: 1811)
- 5'-GG GCCTGCBGGG CC-3' (FRAG.NO:1801) (SEQ. ID NO: 1812) 45
 - 5'-GGCBGCB BGG-3' (FRAG.NO:1802) (SEQ. ID NO: 1813)
 - 5'-GGG TCC TCB TGG CTG GGG-3' (FRAG. NO:1212) (SEQ. ID NO:1222)
 - 5'-GCC TGG GCC TGC BGG GCC-3' (FRAG. NO:1213) (SEQ. ID NO:1223)
 - 5'-GCT CTT GCC TGG BGT GGC TC-3' (FRAG. NO:1214) (SEQ. ID NO:1224)
- 5'-CCGGGGCTGC BGCBBCCTCB TCBGCTCTTG CCTGGBGTGG CTCBGCCTGG GCCTGCBGGG CCBCCBGGBG BBTGGCBGCB GGGBTGGCGB GGGTCCTCBT GGCTGGGGTC BCBGBTCCTC TBGCTBGGCB GGGTGBCCBG BGBGGGC-3' (FRAG.NO:1803) (SEQ. ID NO: 1814) 5'-GCC CBG BGT CTT CCC TGG T-3' (FRAG. NO:1215) (SEQ. ID NO:1225) 50

Human Defensin 3 Anti-sense Oligonucleotide Fragments

5'-CGCTGCBBTC TGCTCCGGGG CTGCBGCBBC CTCBTCBGCTC TTGCCTGGBGTG GCTCBGCCTGG GCCTGCBGGG CCBCCBGGBGB BTGGCBGCBBGG GBTGGCGBGGG TCCTCBTGGC TGGGGTCBCCT GGBGGBGGGB GBGCBGGGG TCCTCBTGGC TGGGGTCCCT CTCTCCCGTC CT-3' (FRAG. NO:1804) (SEQ. ID NO:1815)

5'-GGCBGCBBGG-3' (FRAG. NO:1805) (SEQ. ID NO:1816)

5'-GG CTG GGG-3' (FRAG. NO:1806) (SEQ. ID NO:1817)

5'-GGGGTCBCC-3' (FRAG. NO:1807) (SEQ. ID NO:1818)

5'-GGG TCC TCB TGG CTG GGG TC-3' (FRAG. NO:1216) (SEQ. ID NO:1226)

5'-CCT CTC TCC CGT CCT-3' (FRAG. NO:1217) (SEQ. ID NO:1227)

5'-CGCTGCBBTC TGCTCCGGGG CTGCBGCBBC CTCBTCBGCTC TTGCCTGGBGTG GCTCBGCCTGG GCCTGCBGGG CCBCCBGGBGB BTGGCBBG GBTGGCGBGGG TCCTCBTGGC TGGGGTCBCCT GGBGGBGGGB GBGCBGG-3' 10 NO:1808) (SEQ. ID NO:1819)

Human Macrophage Inflammatory Protein-1-alpha/RANTES Receptor Anti-sense Oligonucleotide Fragments

- 5'-GTCTTTGTTT CTGGGCTCGT GCCCCBTCCC GGCTTCTCTC TGGTTCCGTC CTCTGTGGTG TTTGGCCCTG CTTCCTTTTG GGGCAGCAGT TGGGCCCCAA AGGCCCTCTC GTTCACCTTC TGGCACGGAGTT GCATCCCCATA CCTGTTGAGG GTCAAACTCT GTGGTCGTGT CATAGTCCTC TGTGGTGTTT GGAGTTTCCA TCCCGGCTTC TCTCTGGTTC CAAGGGAGB 15 GGGGGCBGCB GTTGGGCCCC BBBGGCCCTC TCGTTCBCCT TCTGGCBCGG BGTTGCBTCC CCBTBGTCBB BCTCTGTGGT CGTGTCBTBG TCCTCTGTGG TGTTTGGBGT TTCCBTCCCG GCTTCTCTCT GGTTCCBBGG GB-3' (FRAG. NO:1809) (SEQ. ID NO:1820)
- 5'-GGGCC CC-3' (FRAG. NO:1810) (SEQ. ID NO:1821) 20

5'-GGGGGCBGC-3' (FRAG. NO:1811) (SEQ. ID NO:1822)

5'-CCCGGCTTC-3' (FRAG. NO:1812) (SEQ. ID NO:1823)

- 5'-GTC TTT GTT TCT GGG CTC GTG CC-3' (FRAG. NO:1218) (SEQ. ID NO:1228) 5'-CCB TCC CGG CTT CTC TCT GGT TCC-3' (FRAG. NO:1219) (SEQ. ID NO:1229)
- 5'-GTC CTCTGT GGT GTT TGG-3' (FRAG. NO:1220) (SEQ. ID NO:1230) 25

5'-CCC TGC TTC CTT TTG CCT GTT-3' (FRAG. NO:1221) (SEQ. ID NO:1231)

GAGGGGGCAGCAGTTGGGCCCCAAAGGCCCTCTCGTTCACCTTCTGGCACGGAGTTGCATCCCCATAGTCAAACTCTGTGGT CGT-3

(FRAG. NO:1222) (SEQ. ID NO:1232) 30

5'-GTCATAGTCCTCTGTGGTGTTTGGAGTTTCCATCCCGGCTTCTCTCTGGTTCCAAGGGA-3 ' (FRAG. NO:1223) (SEQ. ID

5'-GBGGGGGCBG CBGTTGGGCC CCBBBGGCCC TCTCGTTCBC CTTCTGGCBC GGBGTTGCBT CCCCBTBGTC BBBCTCTGTG

GTCGTG-3' (FRAG, NO:1224) (SEQ. ID NO:1234) 5'-TCBTBGTCCTCTGTGGTGTTTGGBGTTTCCBTCCCGGCTTCTCTCTGGTTCCBBGGGB-3' (FRAG. NO:1225) (SEQ. ID NO:1235) 35

RANTES Antisense Oligonucleotide Fragments

5'-GGGCBCGGGG CBGTGGGCGG GCBBTGTBGG CBBBGCBGCB GGGTGTGGTG TCCGBGGBBT BTGGGGBGGC BGBTGCBGGB GCGCBGBGGG CBGTBGCBBT GBGGBTGBCB GCGBGGCGTG CCGCGGBGBC CTTCBTGGTB CCTGTGGBGB GGCTGTCGGB GGGGGTGTGG TGTCCGCTTG GCGGTTCTTT CGGGTGTTTC TTCTCTGGGT TGGCCTGCTG CTCGTCGTGGT CGCTCCGCTC CCGGGTTCGT CTCGCTCTTC TTCCTTGTCG TGTTCCTCCC TTCCTTGCCT CT-3' (FRAG. NO: 1813) (SEQ.

40 ID NO: 1824)

5'-GGGTTGGC-3' (FRAG. NO: 1814) (SEQ. ID NO: 1825)

5'-CGGGG CBG-3' (FRAG. NO: 1815) (SEQ. 1D NO: 1826)

5'-CCCGGGTTCG-3' (FRAG. NO: 1816) (SEQ. ID NO: 1827) 5'-GGGTGTGGTG-3' (FRAG. NO: 1817) (SEQ. ID NO: 1828) 5'-GGGCBCGGGG CBGTGGGCGG GCBBTGTBGG CBBBGCBGCB GGGTGTGGTG TCCGBGGBBT BTGGGGBGGC BGBTGCBGGB 45 GCGC-3' (FRAG. NO:1226) (SEQ. ID NO:1236)

5'-BGBGGGCBGTBGCBBTGBCBGCGBGGCGTGCCGCGGBGBCCTTCBTGGTBCCTGTGGBGGGGCTGTCGGBGG-3'

(FRAG. NO:1227) (SEQ. ID NO:1237) (FRAG. 50

NO:1228) (FRAG. NO:1229)

55 (SEQ. ID NO:1239)

5'-GGGTGTGCTCCG-3' (FRAG. NO:1230) (SEQ. ID NO:1240)

5'-CTTGGCGGTTCTTTCGGGTG-3' (FRAG. NO:1231) (SEQ. ID NO:1241)

5'-TTTICTTCTCTGGGTTGGC-3' (FRAG. NO:1232) (SEQ. ID NO:1242) 5'-CTGCTGCTCGTCGTCGTC-3' (FRAG. NO:1233) (SEQ. ID NO:1243)

- 5'-GCTCCGCTCCCGGGTTC-3' (FRAG. NO:1234) (SEQ. ID NO:1244) 5'-GTCTCGCTCTGTCGCCC-3' (FRAG. NO:1235) (SEQ. ID NO:1245)
- 5'-CTTCCTTCCTTGTC-3' (FRAG. NO:1236) (SEQ. ID NO:1246)
- 5'-GTGTTCCTCCCTTCCTTGCCTCT-3' (FRAG. NO:1237) (SEQ. ID NO:1247)
- 5'-GGCBCGGGG CBGTGGGCGG GCBBTGTBGG CBBBGCBGCB GGGTGTGGTG TCCGBGGBBT BTGGGGBGGC BGBTGCBGGB 5 GCGCBGBGGG CBGTBGCBBT GBGGBTGBCB GCGBGGCGTG CCGCGGBGBC CTTCBTGGTB CCTGTGGBGB GGCTGTCGGB GG-3' (FRAG. NO:1818) (SEQ. ID NO:1829)

Human Muscarinic Acetylcholine Receptor HM1* Anti-sense Oligonucleotide Fragments

- 5'-GCTGCCCGGC GGGGTGTGCG CTTGGCGCTC CCGTGCTCGG TTCTCTGTCT CCCGGTCCCC CTTGCCTGGC GTCTCGGGCC TTCGTCCTCT TCCTCTTCTT CCTTCCGCTC CGTGGGGGCT GCTTGGTGGG GGCCTGTGCCT CGGGGTCCCG GGGCTTCTGG 10 CCCTTGCCGT TCATGGTGGC TAGGTGGGGC GTTCBTGGTG GCTBGGTGGG GC-3' (FRAG. NO:1819) (SEQ. ID NO: 1830)
 - 5'-GGTGGGGC-3' (FRAG. NO:1820) (SEQ. ID NO: 1831)
 - 5'-GCCCGGCGGGG-3' (FRAG. NO:1821) (SEQ. ID NO: 1832)
- 5'-CGG GGC TTC TGG CCC-3' (FRAG. NO:1822) (SEQ. ID NO: 1833)
- 5'-GTT CBT GGT GGC TBG GTG GGG C-3' (FRAG. NO:1238) (SEQ. ID NO:1248) 15
 - 5'-GCT GCC CGG CGG GGT GTG CGC TTG GC-3' (FRAG. NO:1239) (SEQ. ID NO:1249)
 - 5'-GCT CCC GTG CTC GGT TCT CTG TCT CCC GGT-3' (FRAG. NO:1240) (SEQ. ID NO:1250)

 - 5'-CCC CCT TTG CCT GGC GTC TCG G-3' (FRAG. NO:1241) (SEQ. ID NO:1251)
 5'-GCC TTC GTC CTC TTC CTC TTC CTT CC-3' (FRAG. NO:1242) (SEQ. ID NO:1252)
- 5'-GCT CCG TGG GGG CTG CTT GGT GGG GGC CTG TGC CTC GGG GTC C-3' (FRAG. NO:1243) (SEQ. ID NO:1253) 20
 - 5'-CGG GGC TTC TGG CCC TTG CC-3' (FRAG. NO:1244) (SEQ. ID NO:1254)
 - 5'-GTT CAT GGT GGC TAG GTG GGG C-3' (FRAG. NO: 1245) (SEQ. ID NO:1255)

Human Muscarinic Acetylcholine Receptor HM3* Anti-sense Oligonucleotide Fragments

- 5'-GGG GTG GGT BGG CCG TGT CTG GGGGTT GGC CBT GTT GGT TGC CTCT TGG TGC GCC GGG CGCG TCT TGG CTT 25 AGG CCG TGT CTG GGGGTT GGC CAT GTT GGT TGC CGGG CCC GCG GCT GCA GGG G-3' (FRAG. NO:1823) (SEQ. ID NO-1934\
- 5'-CCC GGG CGG-3' (FRAG. NO:1824) (SEQ. ID NO:1835) 30
 - 5'-G GCG GGG GGG CC-3' (FRAG. NO:1825) (SEQ. ID NO:1836)
 - 5'-CCC GGG CCG CC-3' (FRAG. NO: 1826) (SEQ. ID NO: 1837)
 - 5'-GG CCG TGT-3' (FRAG. NO:1827) (SEQ. ID NO:1838)
- 5'-GGG GTG GGT BGG CCG TGT CTG GGG-3' (FRAG. NO:1246) (SEQ. ID NO:1256)
- 5'-GTT GGC CBT GTT GGT TGC C-3' (FRAG. NO:1247) (SEQ. ID NO:1257) 5'-TCT TGG TGC GCC GGG C-3' (FRAG. NO:1248) (SEQ. ID NO:1258) 35

 - 5'-GCG TCT TGG CTT TCT TCT TCT TCG GGC CCT CGG GCC GGT GCT TGT GG-3' (FRAG. NO:1249) (SEQ. ID NO:1259)

 - 5'-GCG CTG GCG GGG GGG CCT CCT CC-3' (FRAG. NO:1251) (SEQ. ID NO:1261)
- 5'-GCT CTG TGG CTG GGC GTT CCT TGG TGT TCT GGG TGG C-3' (FRAG. NO:1252) (SEQ. ID NO:1262) 40
 - 5'-TGG CGG GCG TGG TGG CCT CTG TGG TGG-3' (FRAG. NO:1253) (SEQ. ID NO:1263)
 - 5'-GGG CCC GCG GCT GCB GGG G-3' (FRAG. NO:1254) (SEQ. ID NO:1264)
 - 5'-TTG CCT GTC TGC TTC GTC-3' (FRAG. NO:1255) (SEQ. ID NO:1265)
- 5'-CTT TGC GCT CCC GGG CCG CC-3' (FRAG. NO:1256) (SEQ. ID NO:1266)
- 5'-GGG GTG GGT AGG CCG TGT CTG GGG-3' (FRAG. NO:1257) (SEQ. ID NO:1267) 45
- 5'-GTT GGC CAT GTT GGT TGC C-3' (FRAG. NO:1258) (SEQ. ID NO:1268) 5'-GGG CCC GCG GCT GCA GGG G-3' (FRAG. NO:1259) (SEQ. ID NO:1269)

Human Fibronectin* Antisense Oligonucleotide Fragments

- TCC CCG CCC CGC CCC GGC CCG GGG GGC GGG GGC GG CTC GCG CCT GGG GTT CCC TCT CCC CCT GTG C GCC TGC CTC TTG CTC TTCTGC GTC CGC TGC CTT CTC CC CTC 50 CTG GTG GTG GTG GTG GTG CCT CTG CCC GTG CTC GCCCTG CCT GGG CTG GCC TCT TCG GGT GTG GCT TTG GGG CTC
- 55 GTG GGC TIT CCC TGG TCC GGT CIT CTC CTT GGG GGT C GCC CTT CTT GGT GGG CTGGCT CGT CTG TCT TTT TCC TTC

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5'-GGCCCGGGC-3' (FRAG. NO:1829) (SEQ. ID NO: 1840)
       5'-GCCGGCGCGGCG-3' (FRAG. NO:1830) (SEQ. ID NO:1841)
5'-GCCTGGCTGGCC-3' (FRAG. NO:1831) (SEQ. ID NO: 1842)
       5'-GGGGG TGGCCG-3' (FRAG. NO:1832) (SEQ. ID NO: 1843)
       5'-GG GGG TGG CCG TTG TGG GCG G-3' (FRAG. NO:1833) (SEQ. ID NO: 1844)
       5'-CGG TTT CCT TTG CGG TC-3' (FRAG. NO:1260)(SEQ. ID NO:1270)
       5'-TTG GCC CGG GCT CCG GGT G-3' (FRAG. NO:1261)(SEQ. ID NO:1271)
       5'-CCC GCC CGC CCG CCC GCC GCC GC-3' (FRAG. NO:1262)(SEQ. ID NO:1272)
10
       5'-CCC GCC GGG CTG TCC CCG CCC CGC CCC-3' (FRAG. NO:1263)(SEQ. ID NO:1273)
       5'-GGC CCG GGG CGC GGG GG-3' (FRAG. NO:1264)(SEQ. ID NO:1274)
       5'-CGG CCC TCC CGC CCC TCT GG-3' (FRAG. NO:1265)(SEQ. ID NO:1275)
       5'-GCC GGC GCG GGC GTC GG-3' (FRAG. NO:1266)(SEQ. ID NO:1276)
       5'-CCG CTC GCG CCT GGG GTT CCC TCT CCT CCC CCT GTG C-3' (FRAG. NO:1267)(SEQ. ID NO:1277)
15
       5'-GCC TGC CTC TTG CTC TTC-3' (FRAG. NO:1268)(SEQ. ID NO:1278)
       5'-TGC GTC CGC TGC CTT CTC CC-3' (FRAG. NO:1269)(SEQ. ID NO:1279)
       5'CTC TCC TCG GCC GTT GCC TGT GC-3' (FRAG. NO:1270)(SEQ. ID NO:1280)
       5'-TGT CCG TCC TGT CGC CCT TCC GTG GTG C-3' (FRAG. NO:1271)(SEQ. ID NO:1281)
        5'-TGT TGT CTC TTC TGC CCT C-3' (FRAG. NO:1272)(SEQ. ID NO:1282)
20
        5'-GGT GTG CTG GTG GTG GTG GTG-3' (FRAG. NO:1273)(SEQ. ID NO:1283)
        5'-CCT CTG CCC GTG CTC GCC-3' (FRAG. NO:1274)(SEQ. ID NO:1284)
        5'-CTG CCT GGG CTG GCC TCT TCG GGT-3' (FRAG. NO:1275)(SEQ. ID NO:1285)
        5'-GTG GCT TTG GGG CTC TCT TGG TTG CCC TTT-3' (FRAG. NO:1276)(SEQ. ID NO:1286)
        5'-CTT CTC GTG GTG CCT CTC CTC CCT GGC TTG GTC GT-3' (FRAG. NO:1277)(SEQ. ID NO:1287)
25
        5'- TGT CTG GGG TGG TGC TCC TCT CCC-3' (FRAG. NO:1278)(SEQ. ID NO:1288)
        5'-TTT CCC TGC TGG CCG TTT GT-3' (FRAG. NO:1279)(SEQ. ID NO:1289)
        5'-CCT GTT TTC TGT CTT CCT CT-3' (FRAG. NO:1280)(SEQ. ID NO:1290)
        5'-TTC CTC CTG TTT CTC CGT-3' (FRAG. NO:1281)(SEQ. ID NO:1291)
        5'-TTG GCT TGC TGC TTG CGG GGC TGT CTC C-3' (FRAG. NO:1282)(SEQ. ID NO:1292)
30
        5'-CTT GCC CCT GTG GGC TTT CCC-3' (FRAG. NO:1283)(SEQ. ID NO:1293)
        5'-TGG TCC GGT CTT CTC CTT GGG GGT C-3' (FRAG. NO:1284)(SEQ. ID NO:1294)
        5'-GCC CTT CTT GGT GGG CTG-3' (FRAG. NO:1285)(SEQ. ID NO:1295)
        5'-GCT CGT CTG TCT TTT TCC TTC C-3' (FRAG. NO:1286)(SEQ. ID NO:1296)
        5'-TGG GGG TGG CCG TTG TGG GCG GTG TGG TCC GCC T-3' (FRAG. NO:1287)(SEQ. ID NO:1297)
 35
        5'-TGC CTC TGC TGG TCT TTC-3' (FRAG. NO:1288)(SEQ. ID NO:1298)
        Human Interleukin-8* Fragments Antisense Oligonucleotide Fragments
        5'-GBTGTTTGTT BCCBBBGCBT CBBGBBTBGC TTTGCTBTCT BBGGBTCBCB TTTBGBCBTB GGBBBBCGCT GTBGGTCBGBB
 40
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BGBTGTGCTT BCCTTCBCBC BGBGCTGCBG BBBTCBGGBBGG CTGCCBBGBGBG CCBCGGCCBGC TTGGBGTCBT GTTTBCBCBC BGTGBGGTGC TCCGGTGGCT TTTTGCTTGT GTGCTCTGCT GTCTCTG TTC CTTCCGGTGG TTTCTTCCTG GCTCTTGTCC TTTCTCTTGG CCCTTGGCCC-3' (FRAG. NO:1834) (SEQ. ID NO:1845) 5'-G CTC CGG-3' (FRAG. NO:1835) (SEQ. ID NO:1846) 5'-CBBGBBTBGC-3' (FRAG. NO:1836) (SEQ. ID NO:1847)

5'-CBCBC BGTGBGGTGC-3' (FRAG. NO:1837) (SEQ. ID NO:1848) 5'-BCCBBBGCBT CBBGBBTBGC-3' (FRAG. NO:1838) (SEQ. ID NO:1849) 45 5'-GCCBBGBGBG CCBCGGCCBGC-3' (FRAG. NO:1839) (SEQ. ID NO:1850) 5'-GTG CTC CGG TGG CTT TTT-3' (FRAG. NO:1289)(SEQ. ID NO:1299) 5'-GCT TGT GTG CTC TGC TGT CTC TG-3' (FRAG. NO:1290)(SEQ. ID NO:1300)

5'-'TTC CTT CCG GTG GTT TCT TCC TGG CTC TTG TCC T-3' (FRAG. NO:1291)(SEQ. ID NO:1301)

5'-TTC TCT TGG CCC TTG GCC C-3' (FRAG. NO:1292)(SEQ. ID NO:1302) 5'-GBTGTTTGTT BCCBBBGCBT CBBGBBTBGC TTTGCTBTCT BBGGBTCBCB TTTBGBCBTB GGBBBBCGCT GTBGGTCBGBB 50 BGBTGTGCTT BCCTTCBCBC BGBGCTGCBG BBBTCBGGBBGG CTGCCBBGBGBG CCBCGGCCBGC TTGGBGTCBT GTTTBCBCBC BGTGBGGTGC TCCGGTGGCT TTTTGCTTGT-3' (FRAG. NO:1840) (SEQ. ID NO:1851)

Human IL-8 Receptor Alpha Antisense Oligonucleotide Fragments

5'-ACAGGGGCTG TAATCTTCATC TGCAGGTGGC ATGCCAGTGA AATTTAGATC ATCAAAATCC CACATCTGTG GATCTGTAAT ATTTGACATG TCCTCTTCAG TTTCAGCAAT GGTTTGATCT AACTGAAGCA CCGGCCAGGB CBGGGGCTGT 55 BBTCTTCBTC TGCBGGTGGC BTGCCBGTGB BBTTTBGBTC BTCBBBBTCC CBCBTCTGTG GBTCTGTBBT BTTTGBCBTG TCCTCTTCBG TITCBGCBB TGGTTTGBTC TBBCTGBBGC BCCGGCCBGG TGGCTCGGTG CTTCTGCCCC TGTTGTTGCG GCGCTCGGTT GGTGTGGCCC CTGTGGTGCT TCGTTTCCCC CTCTTTCTCT TTGTTCGGGG GTTCTTGTGG CGGGCTGCTT 60 GTCTCGTTCC-3'

(FRAG. NO:1841) (SEQ. ID NO:1852)

- 5'-CBGGGGC-3' (FRAG. NO:1842) (SEQ. ID NO:1853)
- 5'-GCBGGTGGC-3' (FRAG. NO:1843) (SEQ. ID NO:1854)
- 5'-GCGGCGCTC-3' (FRAG. NO:1844) (SEQ. ID NO:1855)
- 5'-TGGCTCGGTGCTTCTGCCCC (FRAG. NO:1293)(SEQ. ID NO:1303) 5
 - 5'-TGTTGTTGCGGCGCTC (FRAG. NO:1294)(SEQ. ID NO:1304)
 - 5'-GGTTGGTGTGCCCCTG (FRAG. NO:1295)(SEQ. ID NO:1305) 5'-TGGTGCTTCGTTTCC (FRAG. NO:1296)(SEQ. ID NO:1306)
 - 5'-CCCTCTTTCTCTTTGTTC (FRAG. NO:1297)(SEQ. ID NO:1307)
- 5'-GGGGGTTCTTGTGGC (FRAG. NO:1298)(SEQ. ID NO:1308) 10
 - 5'-GGGCTGCTTGTCTCGTTCC (FRAG. NO:1299)(SEQ. ID NO:1309)
 - 5'-ACAGGGGCTG TAATCTTCATC TGCAGGTGGC ATGCCAGTGA AATTTAGATC ATCAAAATCC CACATCTGTG GATCTGTAAT ATTTGACATG TCCTCTTCAG TTTCAGCAAT GGTTTGATCT AACTGAAGCA CCGGCCAGG-3' (FRAG. NO:1845) (SEO, ID NO:1856)
- 5'-B CBGGGGCTGT BBTCTTCBTC TGCBGGTGGC BTGCCBGTGB BBTTTBGBTC BTCBBBBTCC CBCBTCTGTBBT 15 BTTTGBCBTG TCCTCTTCBG TTTCBGCBB TGGTTTGBTC TBBCTGBBGC BCCGGCCBGG-3' (FRAG. NO:1846) (SEQ. ID

Human GM-CSF Antisense Oligonucleotide Fragments

- 5'-CTTGBGCBGG BBGCTCTGGG GCBGGGBGCT GGCBGGGGCCC BGGGGGGTGG CTTCCTGCBC TGTCCBGBGT GCBCTGTGCC CBGGCTCCGG GCGGTCCBGCCBTGGGTCTG GGGGCTGGG CTGCBGGCTC CGGGCGGGCG GGTGCGGGCT GCGTGCTGGG 20 TGCTGGGGGC TGCCCCGCAG GCCCTGC-3' (FRAG. NO:1847) (SEQ. ID NO: 1858)
 - 5'-GBGCBGG BBG-3' (FRAG. NO:1848) (SEQ. ID NO: 1859)
- 5'-GCCBCBGCBGCBGC-3' (FRAG. NO:1849) (SEQ. ID NO: 1860) 25 5'-GGG TGC GGG C-3' (FRAG. NO:1850) (SEQ. ID NO: 1861)

 - 5'-GGT CCB GCC BTG GGT CTG GG-3' (FRAG. NO:1300)(SEQ. ID NO:1310) 5'-GGC TGG GCT GCB GGC TCC GG-3' (FRAG. NO:1301)(SEQ. ID NO:1311)
 - 5'-GCG GGC GGG TGC GGG CTG CGT GCT GGG-3' (FRAG. NO:1302)(SEQ. 1D NO:1312)
- 5'-GGC TGC CCC GCA GGC CCT GC-3' (FRAG. NO:1303)(SEQ. ID NO:1313) 30 5'-CTTGBGCBGG BBGCTCTGGG GCBGGGBGCT GGCBGGGCCC BGGGGGGTGG CTTCCTGCBC TGTCCBGBGT GCBCTGTGCC CBGGCTCCGG GC-3' (FRAG. NO:1851) (SEQ. ID NO: 1862)

Human Tumor Necrosis Factor α Antisense Oligonucleotide Fragments

- 5'-GCBCCGCCTG GBGCCCTGGG GCCCCCCTGT CTTCTTGGGG BGCGCCTCCT CGGCCBGCTC CBCGTCCCGG BTCBTGCTTT CBGTGCTCBT GGTGTCCTTT CCBGGGGBGB GBGGGGCTGG TCCTCTGCTG TCCTTGCTGG TGCTCBTGGT GTCCTTTCCG 35 CCCTGGGGCC CCCCTGTCTT CTTGGGGCCT CTTCCCTCTG GGGGCCGTCT CTCTCCCTCT CTTGCGTCTC TCTCTTTCTC TCTCTCTT CCCCTTTCCC GCTCTTTCTG TCTCGGTGTC TGGTTTTCTC TCTCCGCTGG CTGCCTGTCT GGCCTGCGCT CTTGGCCTGT GCTGTTCCTC CTCCGGTTCC TGTCCTCTCT GTCTGTCGCC CCCTCTGGGG TCTCCCTCTG GGTGGTGGTC TTGTTGCTTG GGCTGGGCTC CGTGTCTCCB GTGCTCBTGG TGTCCGCTGB GGGBGCGTCT GCTGGCGCTG GTCCTCTGCTGTC 40 CTTGCTGGTG CTCBTGGTGT CCTTTCCGCC CTGGGGCCCC CCTGTCTTCT TGGGGCCTCT TCCCTCTGGG GGCCGTCTC TCTCCCTCTC TTGCGTCTCT CTCTTTCTCT CTCTCTTCTC CCCTTTCCCG CTCTTTCTGT CTCGGTGTCT GGTTTTCTCT
- CTCCGCTGGC TGCCTGTCTG GCCTGCGCTC TTGGCCTGTG CTGTTCCTCC TCCGGTTCCT GTCCTCTG TCTGTCGCCC CCTCTGGGGT CTCCCTCTGG CGTGGTGGTC TTGTTGCTTG GGCTGGGCTC CGTGTCTCCB GTGCTCBTGG TGTCCGCTGB GGGBGCGTCT GCTGGC-3' (FRAG. NO:1852) (SEQ. ID NO:1863) 45
- 5'-GGGGCCCCCC-3' (FRAG. NO:1853) (SEQ. ID NO:1864)
 - 5'- GGG GGC CG TCT-3' (FRAG. NO:1854) (SEQ. ID NO:1865)
 - 5'-CCBGGGGBGB GBGGGGCTGG-3' (FRAG. NO:1855) (SEQ. ID NO:1866)
- GCBCCGCCTGGBGCCCTGGGGCCCCCCTGTCTTCTTGGGGBGCGCCTCCTCGGCCBGCTCCBCGTCCCGGBTCBTGCTTTCBGTGC 50 TCBTGGTGTCCTTTCCBGGGGBGBGBGGGG-3' (FRAG. NO:1304) (SEQ. ID NO:1314) CCT CTT CCC TCT GGG GGC CG TCT CTC TCC CTC TCT TGC GTC TCT C TCT TTC TCT CTC TCT CTT CCC C TTT CCC GCT CTT TCT GTC TC GGT GTC TGG TTT TCT CTC TCC GCT GGC TGC CTG TCT GGC CTG CGC TCT T GGC CTG TGC TGT TCC
- 55 CTG GGC TCC GTG TCT C CBG TGC TCB TGG TGT CC-3' (FRAG. NO:1305) (SEQ. ID NO:1315) 5'-GCT GBG GGB GCG TCT GCT GGC GCT GGT CCT CTG CTG TCC TTG CTG GTG CTC BTG GTG TCC TTT CC GCC CTG GGG CCC CCC TGT CTT CTT GGG G CCT CTT CCC TCT GGG GGC CG TCT CTC TCC CTC TCT TGC GTC TCT C TCT TTC TCT CTC TET CTT CCC C TIT CCC GCT CTT TCT GTC TC GGT GTC TGG TTT TCT CTC TCC GCT GGC TGC CTG TCT GGC CTG CGC

TCT T GGC CTG TGC TGT TCC TCC GGT TCC TGT CCT CTC TGT CTG TC GCC CCC TCT GGG GTC TCC CTC TGG C GTG GTG GTC TTG TTG CTT GGG CTG GGC TCC GTG TCT C CBG TGC TCB TGG TGT CC GCT GBG GGB GCG TCT GCT GGC

(FRAG. NO:1306) (SEQ. ID NO:1316)

- 5'-GCT GGT CCT CTG CTG TCC TTG CTG-3' (FRAG. NO:1655) (SEQ. ID NO:1665) 5
 - 5'-GTG CTC BTG GTG TCC TTT CC-3' (FRAG. NO:1656)(SEQ. ID NO:1666)
 - 5'-GCC CTG GGG CCC CCC TGT CTT CTT GGG G-3' (FRAG. NO:1657)(SEQ. ID NO:1667)
 - 5'-CCT CTT CCC TCT GGG GGC CG-3' (FRAG. NO:1658)(SEQ. ID NO:1668)
 - 5'-TCT CTC TCC CTC TCT TGC GTC TCT C-3' (FRAG. NO:1659)(SEQ. ID NO:1669)
- 5'-TCT TTC TCT CTC TCT CTT CCC C-3' (FRAG. NO:1660)(SEQ. ID NO:1670) 10
 - 5'-TTT CCC GCT CTT TCT GTC TC-3' (FRAG. NO:1661)(SEQ. ID NO:1671)
 - 5'-GGT GTC TGG TIT TCT CTC TCC-3' (FRAG. NO:1662)(SEQ. ID NO:1672)
 - 5'-GCT GGC TGC CTG TCT GGC CTG CGC TCT T-3' (FRAG. NO:1663)(SEQ. ID NO:1673)
 - 5'-GGC CTG TGC TGT TCC TCC-3' (FRAG. NO:1664)(SEQ. ID NO:1673)
- 5'-TCC GGT TCC TGT CTC TGT CTG TC-3' (FRAG. NO:1665)(SEQ. ID NO:1675)
 5'-GCC CCC TCT GGG GTC TCC CTC TGG C-3' (FRAG. NO:1666)(SEQ. ID NO:1676) 15
 - 5'-GTG GTG GTC TTG TTG CTT-3' (FRAG. NO:1667)(SEQ. ID NO:1677)
 - 5'-GGG CTG GGC TCC GTG TCT C-3' (FRAG. NO:1668)(SEQ. ID NO:1678)
 - 5'-CBG TGC TCB TGG TGT CC-3' (FRAG. NO:1669)(SEQ. ID NO:1679)
- 5'-GCT GBG GGB GCG TCT GCT GGC-3' (FRAG. NO:1670)(SEQ. ID NO:1680) 20

Human Leukotriene C4 Synthase Antisense Oligonucleotide Fragments

5'-CTCGGTBGBC GCGCTCGBBC TCGGGTGGGC CGGTGGTGBG CGGCGGCGBCB CGCGGBBGGC CCTGCGCGCCC GBGBTCBCCTG CBGGGBGBBG TBGGCTTGCB GCBGGBGCTCC CBGGBGGGTG BCBGCBGCCB GTBGBGCTBC CTCGTCCTTC BTGGTBCCGT CGGTGTGGTG GCBCGGGCTG TGTGTGBBGG CGBGCTGGGC CCCGTCTGCT GCTCCTCGTC

- CTTCA TGG TA CCGTCGGTGT GGTGGCCTCG GGTGGGCCGG TGGTGGGCCG CGCGCGCTCG CGTGGCTCCG GCTCTTCTTT 25 CCCGGCTCCGT CGGCCCGGGG GCCTTGGTCT CCCTCGTCCT TCBTGGTBCC G-3' (FRAG. NO:1856) (SEQ ID NO: 1867)
 - 5'-GCB GCBGGBC-3' (FRAG. NO:1857) (SEQ ID NO: 1868)
 - 5'-CCCGGCTCCG-3' (FRAG. NO:1858) (SEQ ID NO: 1869) 5'-CGGCCCGGGG GCC-3' (FRAG. NO:1859) (SEQ ID NO: 1870)
- 5'-CB CGCGG-3' (FRAG. NO:1860) (SEQ ID NO: 1871) 30
 - 5'-GCC CCG TCT GCT GCT CCT CGT GCC G-3' (FRAG. NO:1307)(SEQ. ID NO:1317)
 - 5'-CCT CGT CCT TCA TGG TAC CGT CGG TGT GGT GGC-3' (FRAG. NO:1308)(SEQ. ID NO:1318)
 - 5'-CTC GGG TGG GCC GGT GGT G-3' (FRAG. NO:1309)(SEQ. ID NO:1319)
 - 5'-GGG CGC GCG CGC TCG CGT-3' (FRAG. NO:1310)(SEQ. ID NO:1320)
- 5'-GGC TCC GGC TCT TCT TCC CCG GCT CCG TCG GCC CGG GGG CCT TGG TCT C-3' (FRAG. NO:1311) (SEQ. ID NO:1321) 35
 - 5'-CCT CGT CCT TCB TGG TBC CG-3' (FRAG. NO:1312)(SEQ. ID NO:1322)
 - 5'-CTCGGTBGBC GCGCTCGBBC TCGGGTGGGC CGGTGGTGBG CGGCGGCGBCB CGCGGBBGGC CCTGCGCGCC GBGBTCBCCTG CBGGGBGBBG TBGGCTTGCB GCBGGBCTCC CBGGBGGGTG BCBGCBGCCB GTBGBGCTBC CTCGTCCTTC BTGGTBCCGT CGGTGTGGTG GCBCGGGCTG TGTGTGBBGG CGBGCTGG-3' (FRAG. NO:1861) (SEQ ID NO: 1872)

Human Endothelin-1 Antisense Oligonucleotide Fragments 40

- 5'-BCCGGCGGBG CCGCCBGGGT GGBCTGGGBG TGGGTTTCTC CCCGCCGTTC TCBCCCBCCG CGCTGBGCTC BGCGCCTBBG BCTGCTGTTT CTGGBBGCTCC TTGGCBBGCC BCBBBCBGCB GBGBGBBBBT CBTGBGCBBB TBBTCCBTTC TGBBBBBBBBG GGBTCBBBBB CCTCCCGTTC CCCGTTCGCC TGGCGCGCCG TGCGGGTTCC TCGTGGGTTT CTCCCCGCCG TTCTCCGGTC TGTTGCCTTT GTGGGCTTCT TGTCTTTTTG GCTGTTCTTT TCCTGCTTGG CGTCTTTTCC TTTCTTTGTG CTCGGTTGTG
- GGTCCGCTGG TCCTTTGCCC TGTGTGTTTC TGCTGCCCGT TCGCCTGGCG CGCGCTGCGG GTTCCTCGTG GGTTTCTCCC 45 CGCCGTTCTC CGGTCTGTTG CCTTTGTGGG CTTCTTGTCT TTTTGGCTGT TCTTTTCCTG CTTGGCGTCT TTTCCTTTCT TTGTGCTCGG TTGTGGGTCC GCTGGTCCTT TGCCCTGTGT GTTTCTGCTG-3' (FRAG. NO:1862) (SEQ. ID NO:1873) 5'-CCGGCGGBG CCGCCBGGGT GGBC-3' (FRAG. NO:1863) (SEQ. ID NO:1874)
- 5'-CCGCCBGGG-3' (FRAG. NO:1864) (SEQ. ID NO:1875) 5'-GGCGCGCGC-3' (FRAG. NO:1865) (SEQ. ID NO:1876)
- 50
 - 5'-GTGGGTCCGC-3' (FRAG. NO:1866) (SEQ. ID NO:1877)
 - 5'-CCCGTTCGCCTGGCGC-3' (FRAG. NO:1313)(SEQ. ID NO:1323) 5'-GCGCTGCGGGTTCCTC-3' (FRAG. NO:1314)(SEQ. ID NO:1324)
 - 5'-GTGGGTTTCTCCCCGCCGTTCTC-3' (FRAG. NO:1315)(SEQ. ID NO:1325)
- 5'-CGGTCTGTTGCCTTTGTGGG -3' (FRAG. NO:1316)(SEQ. ID NO:1326) 55
 - 5'-CTTCTTGTCTTTTTGGCT-3' (FRAG. NO:1317)(SEQ. ID NO:1327) 5'-GTTCTTTTCCTGCTTGGC-3' (FRAG. NO:1318)(SEQ. ID NO:1328)

 - 5'-GTCTTTTCCTTTCTT-3' (FRAG. NO:1319)(SEQ. ID NO:1329)
 - 5'-TGTGCTCGGTTGTGGGTC-3' (FRAG. NO:1320)(SEQ. ID NO:1330)
- 5'-CGCTGGTCCTTTGCC-3' (FRAG. NO:1321)(SEQ. ID NO:1331) 60

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5'-CTGTGTGTTTCTGCTG-3' (FRAG. NO:1322)(SEQ. ID NO:1332)
        5'-CCCGTTCGCCTGGCGC-3' (FRAG. NO:1323)(SEQ. ID NO:1333)
5'-GCGCTGCGGGTTCCTC-3' (FRAG. NO:1324)(SEQ. ID NO:1334)
        5'-GTGGGTTTCTCCCCGCCGTTCTC-3' (FRAG. NO:1325(SEQ. ID NO:1335)
        5'-CGGTCTGTTGCCTTTGTGGG-3' (FRAG. NO:1326)(SEQ. ID NO:1336)
        5'-CTTCTTGTCTTTTTGGCT-3' (FRAG. NO:1327)(SEQ. ID NO:1337)
5'-GTTCTTTTCCTGCTTGGC-3' (FRAG. NO:1328)(SEQ. ID NO:1338)
        5'-GTCTTTTCCTTTCTT-3' (FRAG. NO:1329)(SEQ. ID NO:1339)
        5'-TGTGCTCGGTTGTGGGTC-3' (FRAG. NO:1330)(SEQ. ID NO:1340)
         5'-CGCTGGTCCTTTGCC-3' (FRAG. NO:1331)(SEQ. ID NO:1341)
10
         5'-CTGTGTGTTTCTGCTG-3' (FRAG. NO:1332)(SEQ. ID NO:1342)
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Endothelin Receptor ET-B Antisense Oligonucleotide Fragments

5'-GCCCTGTCGG GCGGGAAGCC TCTCTCCTCT CCCCAGATC CGCGACAGGC CGCAGGCAAG AACCAGCGCA ACCAGGGCGC GTCCGCACAG ACTTGGAGGC GGCTGCATGC TGCTACCTGC TCCAGAAGCG TCCGGTGGCC GCCGCGCC CTGTCGGGCG GGBBGCCTCT CTCCTCTCCC CBGBTCCGCG BCBGGCCGCB GGCBBGBBCC BGCGCBBCCB GGGCGCGTCC GCBCBGBCTT GGBGGCGGCT GCBTGCTGCT BCCTGCTCGGGCG GGBBGCCTCCG GTGGCCGCCG CGCGTCCGGT GGCCGCCGCG 15 CCTCTCTCCT CTCCCCGTGG CCCTGTCGGG CGGGTCCTGC CGTCCTGTCT CCTTTTCTTT TGCTGTCTTG TCTTCCCGTC TCTGCTTT-3'

(FRAG. NO: 1867) (SEQ. ID NO: 1878)

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5'-CGGGCG GGBBGCC-3' (FRAG. NO: 1868) (SEQ. ID NO: 1879) 20

5'-CGGGCGGG-3' (FRAG. NO: 1869) (SEQ. ID NO: 1880)

5'-CCGCBCBGBC-3' (FRAG. NO: 1870) (SEQ. ID NO: 1881)

5'-GCGTCCGGTGGCCGCCGC-3' (FRAG. NO:1333)(SEQ. ID NO:1343)

5'-GCCTCTCTCCCCC-3' (FRAG. NO:1334)(SEQ. ID NO:1344)

5'-GTGGCCCTGTCGGGCGGG-3' (FRAG. NO:1335)(SEQ. ID NO:1345)

5'-TCCTGCCGTCCTGTCTCCTTT-3' (FRAG. NO:1336)(SEQ. ID NO:1346)

5'-TCTTTTGCTGTCTTGT-3' (FRAG. NO:1337)(SEQ. ID NO:1347)

5'-CTTCCCGTCTCTGCTTT-3' (FRAG. NO:1338)(SEQ. ID NO:1348)

5'-GCCCTGTCGG GCGGGAAGCC TCTCTCCTCT CCCCAGATC CGCGACAGGC CGCAGGCAAG AACCAGCGCA ACCAGGGCGC

GTCCGCACAG ACTTGGAGGC GGCTGCATGC TGCTACCTGC TCCAGAAGCG TCCGGTGGCC GCCGC-3' (FRAG. NO: 1871) (SEO, ID NO: 1882)

5'-GCCTGTCGG GCGGGBBGCC TCTCTCCTCT CCCCBGBTCC GCGBCBGGCC GCBGGCBBGB BCCBGCGCB BCCBGGGCGC GTCCGCBCBG BCTTGGBGGC GGCTGCBTGC TGCTBCCTGC TCCBGBBGCG TCCGGTGGCC GCCGC-3' (FRAG. NO: 1872) (SEQ. ID NO: 1883)

Endothelin ETA Receptor Antisense Oligonucleotide Fragments 35

5'-GTCTGTCCTC CCCGTCTCCT CCCACTGCTT CTCCCGGGGG CTTCCCCGGC TTCGGGTGGC CGGTGTCCCG GGCTCCGGCG CGGCGGCGGC TTCGCCTGCG GGTGGGTGGC GCGGGCTGCC GGGTCCGCGC GGCGCCTGGG CCCTTGTGCT GCTTTTTGCT TGTTCCGTTC TGGCTGCTCC GGTCTGTGTT GTGGTTGTTT TGTTTCTTCT TGGGTGTGGG CCTTGCGGTT TTGGCTGTGG GCCCTTTGGG GCCTTGGCTT CTGGCTCGTC TGTCCTCCCC GTCTCCTCCC ACTGCTTCT CCCGGGGGCT TCCCCGGCTT CGGGTGGCCG GTGTCCCGGG CTCCGGCGCG GCGGCGGCTT CGGCTGCGGG TGGGTGGCGC GGGCTGCCGG GTCCGCGCGG 40 CGCCTGGGCC CTTGTGCTGC TTTTTGCTTG TTCCGTTCTG GCTGCTCCGG TCTGTGTTGT GGTTGTTTTG TTTCTTCTTG GGTGTGGGCC TTGCGGTTTT GGCTGTGGGC CCTTTGGGGC CTTGGCTTCT GGCTCCAT CCACATGATT GCTTAGATTT GTGCTGTATC TCTCAGGATT ATCACTGATT ACACATCCAA CCAGTGCCAG CCAAAAGGAT GCCCTGAGGC AAAGGGTTTC CATCTTGAGG CAAATTTGAG GACBTCCBC BTGBTTGCTT BGBTTTGTGC TGTBTCTCTC BGGBTTBTCB CTGBTTBCBC BTCCBBCCBG TGCCBGCCBB BBGGBTGCCC TGBGGCBBBG GGTTTCCBTC TTGBGGCBBB TITGBGGB-3' (FRAG. NO:1873) 45

(SEQ. ID NO: 1884)

5'-GBGGCBBBGGG-3' (FRAG. NO:1874) (SEQ. ID NO: 1885)

5'-GCCBGCCBB BBGGB-3' (FRAG. NO:1875) (SEQ. ID NO: 1886)

5'-CGCCTGGGCC C-3' (FRAG. NO:1876) (SEQ. ID NO: 1887)

5'-GTCTGTCCTCCCGGTCTCCCC-3' (FRAG. NO:1339)(SEQ. ID NO:1349) 5'-ACTGCTTCTCCCGGGG-3' (FRAG. NO:1340)(SEQ. ID NO:1350) 50

5'-GCTTCCCCGGCTTC-3' (FRAG. NO:1341)(SEQ. ID NO:1351)

5'-GGGTGGCCGGTGTCCCGGGCTCCGGCGCGCGCGCGC-3' (FRAG. NO:1342)(SEQ. ID NO:1352) 5'-GGCTTCGGCTGC-3' (FRAG. NO:1343)(SEQ. ID NO:1353)

5'-GGGTGGGTGGCGCGG-3' (FRAG. NO:1344)(SEQ. ID NO:1354) 55

5'-GCTGCCGGGTCCGCGCGCGCCCTGGGCC-3' (FRAG. NO:1345)(SEQ. ID NO:1355)

5'-CTTGTGCTGCTTTT-3' (FRAG. NO:1346(SEQ. ID NO:1356)

5'-TGCTTGTTCCGTTC-3' (FRAG. NO:1347)(SEQ. ID NO:1357)

5'-TGGCTGCTCCGGTCTGTTGTGGTTGTTTTG-3' (FRAG. NO:1348)(SEQ. ID NO:1358)

- 5'-TTTCTTCTTGGGTGTGGG-3' (FRAG. NO:1349)(SEQ. ID NO:1359)
- 5'-CCTTGCGGTTTTGG-3' (FRAG. NO:1350)(SEQ. ID NO:1360)
- 5'-CTGTGGGCCCTTTG-3' (FRAG. NO:1351)(SEQ. ID NO:1361)
- 5'-GGGCCTTGGCTTCTGGCTC-3' (FRAG. NO:1352)(SEQ. ID NO:1362)
- 5'-CATCCACATG ATTGCTTAGA TTTGTGCTGT ATCTCTCAGG ATTATCACTG ATTACACATC CAACCAGTGC CAGCCAAAAG 5 GATGCCCTGA GGCAAAGGGT TTCCATCTTG AGGCAAATTT GAGGA-3' (FRAG. NO:1353)(SEQ. ID NO:1363) 5'-CBTCCBCBTG BTTGCTTBGB TTTGTGCTGT BTCTCTCBGG BTTBTCBCTG BTTBCBCBTC CBBCCBGTGC CBGCCBBBBG GBTGCCCTGB GGCBBBGGGT TTCCBTCTTG BGGCBBBTTT GBGGB-3' (FRAG. NO:1354)(SEQ. ID NO:1364)

Substance P Antisense Oligonucleotide Antisense Oligonucleotide Fragments

- 5'-CTGCTGBGGC TTGGGTCTCC GGGCGBTTCT CTGCBGBBGB TGCTCBBBGG GCTCCGGCBG TTCCTCCTTG BTCTGGTCGCT 10 GTCGTBCCBG TCGGBCCBGT BBTTCBGBTC BTCBTTGGCT CCTBTTTCTT CTGCBBBCBG CTGBGTGGBG BCBBGBBBBB BGBCTGCCBB GGCCBCGBGG BTTTTCBTGT TGGBTTTTGC GBCGGBCBGT CCCGCGGGGT GCTGAGTTTC TCTGGTTCCT CCGBGCGCBC GTGGTCGCTC CGCGTTTCTC TGGTTCCTCC GGTCCCGCGG GGTGCTGTCT GGTCGCTGTC GTGGCTTGGG TCTCCGGGCG GTTTCCTTCC TTTTCCGC-3' (FRAG. NO:1877) (SEQ ID NO: 1888)
- 5'-CTCC GGGCGB-3' (FRAG. NO:1878) (SEQ ID NO: 1889) 15 5'-GGCCBCGBGG-3' (FRAG. NO:1879) (SEQ ID NO: 1890)
 - 5'-GGGTCTCCGGGCG-3' (FRAG. NO:1880) (SEQ ID NO: 1891)
 - 5'-GGG TCTCCGGGCG G-3' (FRAG. NO:1881) (SEQ ID NO:1892)
 - 5'-CGTGGTCGCTCCGC-3' (FRAG. NO:1355)(SEQ. ID NO:1365)
- 5'-GTTTCTCTGGTTCCTCCG-3' (FRAG. NO:1356)(SEQ. ID NO:1366) 20
 - 5'-GTCCCGCGGGGTGCTG-3' (FRAG. NO:1357)(SEQ. ID NO:1367) 5'-TCTGGTCGCTGTCGT-3' (FRAG. NO:1358)(SEQ. ID NO:1368)
 - 5'-GGCTTGGGTCTCCGGGCG-3' (FRAG. NO:1359)(SEQ. ID NO:1369)
 - 5'-GTTTCCTTCCTTTTCCGC-3' (FRAG. NO:1360)(SEQ. ID NO:1370)
- 5'-CTGCTGBGGC TTGGGTCTCC GGGCGBTTCT CTGCBGBBGB TGCTCCBBBGG GCTCCGGCBG TTCCTCCTTG BTCTGGTCGCT 25 GTCGTBCCBG TCGGBCCBGT BBTTCBGBTC BTCBTTGGCT CCTBTTTCTT CTGCBBBCBG CTGBGTGGBG BCBBGBBBBB BGBCTGCCBB GGCCBCGBGG BTTTTCBTGT TGGBTTTTGC GBCGGBCBGT CCCGCGGGGT GCTGAGTTTC TCTGGTTCCT CCGBGCGCB-3' (FRAG. NO:1882) (SEQ ID NO: 1893)

Substance P Receptor Antisense Oligonucleotide Fragments

- 5'-GGGCTBBGBT GBTCCBCBTC BCTBCCBCGT TGCCCBCCBC BGBGGTCBCC BCBBTGBCCG TGTBGGCBGC TGCCCBBBGG 30 BCBBTTTGCC BGGCTGGTTG CBCGBBCTGB TTGGGTTCCG BGGTGTTBGT GGBGBTGTTT GGGGBGBGGT CTGBGTCCBC CGGGBGGBCG TTBTCCBTTT CGBBGCTBGG CGGTBBBGCC CTBCTBTCTG TBCBCBBCCC CCCTCTGCBG CBGBGTCCTG TCGTGGCGCC TGGGGCTCBG GGTCCGGGC TAAGATGATC CACATCACTA CCACGTTGCC CACCACAGAG GTCACCACAA TGACCGTGTA GGCAGCTGCC CAAAGGACAA TTTGCCAGGC TGGTTGCACG AACTGATTGG GTTCCGAGGT GTTAGTGGAG
- ATGTTTGGGG AGAGGTCTGA GTCCACCGGG AGGACGTTAT CCATTTCGAA GCTAGGCGGT AAAGCCCTAC TATCTGTACA 35 CAACCCCCCT CTGCAGCAGA GTCCTGTCGT GGCGCCTGGG GCTCAGGGTC CGTCCTGTCG TGGCGCCTGG GGCTCTTCTT TTGTGGGCTC TTTGGTGGCT GTGGCTGTGG TCTCTGTGGT TGCTGCCCTG GGTCTGGGGG TGTGGCCTTG GGGCCGTCCT CTGGCTCCTC CTCGTGGGCC CCC-3' (FRAG. NO:1883) (SEQ. ID NO:1894)
- 5'-GGGBGGBCG-3' (FRAG. NO:1884) (SEQ. ID NO:1895)
- 5'-GGGTC CG-3' (FRAG. NO:1885) (SEQ. ID NO:1896) 40
 - 5-'GGGCC CCC-3' (FRAG. NO:1886) (SEQ. ID NO:1897)
 - 5'-GTCCTGTCGTGGCGCCTGGGGCTC-3' (FRAG. NO:1361)(SEQ. ID NO:1371)
 - 5'-TTCTTTTGTGGGCT-3' (FRAG. NO:1362)(SEQ. ID NO:1372)
 - 5'-CTTTGGTGGCTGTGGCTG-3' (FRAG. NO:1363)(SEQ. ID NO:1373)
- 5'-TGGTCTCTGTGGTTG-3' (FRAG. NO:1364)(SEQ. ID NO:1374) 45
 - 5'-CTGCCCTGGGTCTGG-3' (FRAG. NO:1365)(SEQ. ID NO:1375)
 - 5'-GGGTGTGGCCTTGGGCCCTCCTCGTGGGCCCCC (FRAG. NO:1366)(SEQ. ID NO:1376)
 - 5'-GGGCTAAGAT GATCCACATC ACTACCACGT TGCCCACCAC AGAGGTCACC ACAATGACCG TGTAGGCAGC TGCCCAAAGG ACAATTTGCC AGGCTGGTTG CACGAACTGA TTGGGTTCCG AGGTGTTAGT GGAGATGTTT GGGGAGAGGT CTGAGTCCAC CGGGAGGACG TTATCCATTTC GAAGCTAGGC GGTAAAGCCC TACTATCTGTA CACAACCCCC CTCTGCAGCA GAGTCCTGTC
- 50 GTGGCGCCTG GGGCTCAGGGTCC-3' (FRAG. NO:1367)(SEQ. ID NO:1377) 5'-GGGCTBBGBT GBTCCBCBTC BCTBCCBCGT TGCCCBCCBC BGBGGTCBCC BCBBTGBCCG TGTBGGCBGC TGCCCBBBGG BCBBTTTGCC BGGCTGGTTG CBCGBBCTGB TTGGGTTCCG BGGTGTTBGT GGBGBTGTTT GGGGBGBGGTC TGBGTCCBCC GGGBGGBCGT TBTCCBTTTC GBBGCTBGGC GGTBBBGCCC TBCTBTCTGTB CBCBBCCCCC CTCTGCBGCB GBGTCCTGTC
- GTGGCGCCTG GGGCTCBGGG TCC-3' (FRAG. NO:1368) (SEQ. ID NO:1378) 55

Chymase Antisense Oligonucleotides Antisense Oligonucleotide Fragments

5'-GGBGCTGBTB CTGCBGATTT CBGBGGGBBG BBCCCTGBTB CTCBCCBGCT TCBGCTCTGG BGCBCBBGBG BBBGBGCBGC BGGGGGBGBG GBBGBBGCBG CBTCTTCCCB GBGBGGCTGC CTGBGCBBBT GCTGGTTTTC CTTTCCBGTC TTGGGTTTTB

GGTGGGGTGG CTTCCTTGTT CCTGGGGGTG TCCTCTTGCT CTGGGCTTTT CTCCCCTTTT CCTTCCTGTC TGTTTTCCTG GGGCTCTCCT CTGTCTCTGT GTCCTTGCCC TGGCCCTCTT CCCTCTCTG TCTCCTGTCC CTGTGTTCCG CCCGTCTTCC CTCTCCTGAC CTCCTTTTCC TCCGCTGGGT GGGGCCCTGC CTGTTCTCTG CTCCCTGGCT TGGGGTTTCT TCTGTGTGTC TTCTTCCTCT GTTGGCTGGC TTTCTCCTTC TTTTGTCTTC CTGGGTGCCC CTTCTTCCTT TCTTGGGTCC TTGGTGCTTG GGCTGGG-3' (FRAG. NO:1887) (SEQ. ID NO:1898) 5'-GGBGCBCBBG-3' (FRAG. NO:1888) (SEQ. ID NO:1899) 5'-GBBGCBGC-3' (FRAG. NO:1889) (SEQ. ID NO:1900) 5'-GGGGCBBGG CG-3' (FRAG. NO:1890) (SEQ. ID NO:1901) 5'-CGTTTTCTCTCTC-3' (FRAG. NO:1369)(SEQ. ID NO:1379) 10 5'-GCTGGTTTTCCTTTCC-3' (FRAG. NO:1370)(SEQ. ID NO:1380) 5'-TTCCTTGTTCCTGGGGGTGTCCT-3' (FRAG. NO:1372)(SEQ. ID NO:1382) 5'-CTTGCTCTGGGCTTTTCT-3' (FRAG. NO:1373)(SEQ. ID NO:1383) 5'-CCCCTTTTCCTTCC-3' (FRAG, NO:1374)(SEQ, ID NO:1384) 15 5'-TGTCTGTTTTCCTGGGG-3' (FRAG. NO:1375)(SEQ. ID NO:1385) 5'-CTCTCCTCTGTCTCTGTGT-3' (FRAG. NO:1376)(SEQ. ID NO:1386) 5'-CCTTGCCCTGGCCC-3' (FRAG. NO:1377)(SEQ. ID NO:1387) 5'-TCTTCCCTCTCCTGTCTCCTGT-3' (FRAG. NO:1378)(SEQ. ID NO:1388) 5'-CCCTGTGTTCCGCCC-3' (FRAG. NO:1379)(SEQ. ID NO:1389) 20 5'-GTCTTCCCTCTCTG-3' (FRAG. NO:1380)(SEQ. ID NO:1390) 5'-ACCTCCTTTTCCTCCG-3' (FRAG. NO:1381)(SEQ. ID NO:1391) 5'-CTGGGTGGGCCCTG-3' (FRAG. NO:1382)(SEQ. ID NO:1392) 5'-CCTGTTCTCTGCTCCC-3' (FRAG. NO:1383)(SEQ. ID NO:1393) 5'-TGGCTTGGGGTTTCTTCTG-3' (FRAG. NO:1384)(SEQ. ID NO:1394) 25 5'-TGTGTCTTCTTCTCTGTT-3' (FRAG. NO:1385)(SEQ. ID NO:1395) 5'-GGCTGGCTTTCTCCTTC-3' (FRAG. NO:1386)(SEQ. ID NO:1396) 5'-TTTTGTCTTCCTGGG-3' (FRAG. NO:1387)(SEQ. ID NO:1397) 5'-TGCCCCTTCTTCCTTTCTTGGG-3' (FRAG. NO:1388)(SEQ. ID NO:1398) 5'-TCCTTGGTGCTTGGGCTGGG-3' (FRAG. NO:1389)(SEQ. ID NO:1399) 5'-GGBGCTGBTB CTGCBGATTT CBGBGGGBBG BBCCCTGBTB CTCBCCBGCT TCBGCTCTGG BGCBCBBGBG BBBGBGCBGC 30 BGGGGBBG GBBGBBGCBG CBTCTTCCCB GBGBGGCTGC CTGBGCBBBT GCTGGTTTTC CTTTCCBGTC TTGGGTTTTB

TBBCTCCCBG BBGGCBBGBG BGGGGCBBGG-3' (FRAG. NO:1891) (SEQ. ID NO:1902)

- Endothelial Nitric Oxide Synthase Antisense Oligonucleotide Fragments 5'-GCGTCTTGGG GTGCBGGGCC CBTCCTGCTG CGCCTGGGCG CTGCTGTGCG TCCGTCTGCT GGGGGGCCGG GGTGGCTGGG 35 CCCTGCTTGC CGCACGACCC CGGGCCGACC CGAGGCTCGG GGGGCTGTGT TCTGGCGCTG GTGGGCTTGG GCCCCTCTGG GGGCTGGGTT TCCTGCTGCG CCTGGGCGCT GGCGTCTTGG GGTGCGGGGC CGGGGGGCCG GGGGGCCGCT GTTCGTGGGC TTTCCTGGGG TCCGCGTGGG GTGCTCCGGT TCCTCGTGCC GCTGCTGCCT TGTCTTTCCG GCCGTGGCGG CGTGGTGGTC CGCCCCCCT GGCCTTCTGC TCGGGGTCTG GCTGGTTGCC GGTGCCCTTG GCGGCGGTCT TCTTCCTGGT GGCTCTGGGC 40 CCGGCCGGTC TCGGGCGTCT CGTGTTCGCT CTTGTGCTGT TCCGGCCGCT CCTTCCTCTT CCGCCGCCGC CGCTCCCCGC CCGCTCGTCG CCCTGGCCCG GCCTCCTCCT GGCCGCTGTC TCGGGCGGCG GCCTTGGCGC TCCGTTTGGG GCTGCCTCTG GCGCTTCCGG CCCTCGGCCT GGGCGCTCTC TTCCGCCTGT GCTGGTGGCC CTCGTGGGCC CCTCCTGGCC TCCGGTGTCC TGTGGTCCCC CGGCTGGTGG CCGGCCGGT TGGGCGGCGG TGGGCGCCGG CGGGTCCTCC GGGCTGCCCT TCTCCGCCGG GGGTCCCGCG CTCCTGCTGT TCCCTGGGCT CTTCTGCCTC TCTCCTGGGT GGGTGCTGGG TGCCGGGGTC TCCGGGCTTG 45 CCCCGCGCTG CTGGGCGTTC TGCGGTCTTG GGGTTGTCTG TGGCCCCGCT CGTGTCGCCC TCCGTCGCCC GTCGCCGGCC TCGTCCCCTC CTGGGTGCGC GGCGGGCTGG TCCTGGCGTT TTGCTCCTTC CTGGGCGTCT TGGGGTGCBG GGCCCBTCCT GCTGCGCCTG GGCGCTGCTG TGCGTCCGTC TGCTGGGGGG CCGGGGTGGC TGGGCCCTGC TTGCCGCACG ACCCCGGGCC GACCCGAGGC TCGGGGGGCT GTGTTCTGGC GCTGGTGGGC TTGGGCCCCT CTGGGGGCTG GGTTTCCTGC TGCGCCTGGG CGCTGGCGTC TTGGGGTGCG GGGCCGGGGG GCCGGGGGGC CGCTGTTCGT GGGCCTGGGG GTGCCTGTGG CTGCCGGTTG 50 CCCCGGTTGG TGGCGCCGTC CTGCTGCCGG TCGTTGGCTG GGTCCCCCCG CCCGTTTCCT GGGGTCCGCG TGGGGTGCTC CGGTTCCTCG TGCCGCTGCT GCCTTGTCTT TCCGGCCGTG GCGGCGTGGT GGTCCGCCCC CCCTGGCCTT CTGCTCGGGG TCTGGCTGGT TGCCGGTGCC CTTGGCGGCG GTCTTCTTCC TGGTGGCTCT GGGCCCGGCC GGTCTCGGGC GTCTCGTGTT TCCTGGCCGC TGTCTCGGGC GGCGGCCTTG GCGCTCCGTT TGGGGCTGCC TCTGGCGCTT CCGGCCCTCG GCCTGGGCGC TCTCTTCCGC CTGTGCTGGT GGCCCTCGTG GGCCCCTCCT GGCCTCCGGT GTCCTGTGGT CCCCCGGCTG GTGGCCGGGC 55 CGGTTGGGCG GGCGTGGGCG CCGGGGGGTC CTCCGGGCTG CCCTTCTCCG CCGGGGGTCC CGCGCTCCTG CTGTTCCCTG GGCTCTTCTG CCTCTCTCCT GGGTGGGTGC TGGGTGCCGG GGTCTCCGGG CTTGCCCCGC GCTGCTGGGC GTTCTGCGGT CTIGGGGTTG TCTGTGGCCC CGCTCGTGTC GCCCTCCGTC GCCCGTCGCC GGCCTCGTCC CCTCCTGGGT GCGCGGCGGG CTGGTCCTGG CGTTTTGCTC CTTCCTGG-3' (FRAG. NO:1892) (SEQ. ID NO: 1903) 60 5'-GCGGGGCCG-3' (FRAG. NO:1893) (SEQ. ID NO: 1904)
 - 5'-CGGGGGGC-3' (FRAG. NO:1894) (SEQ. ID NO: 1905)

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5'-GCGCGGCGGC-3' (FRAG. NO:1895) (SEQ. ID NO: 1906) 5"-CTGTGCGTCCGTCTGCTGG (FRAG. NO:1390)(SEQ. ID NO:1400) GGGGCCGGGGTGGCCGGCCTGCTTGCCGC (FRAG. NO:1391)(SEQ. ID NO:1401) ACGACCCCGGGCCGACCCGAG (FRAG. NO:1392)(SEQ. ID NO:1402) GCTCGGGGGGCTGTTCTGGCGCTGGTGGG (FRAG. NO:1393)(SEQ. ID NO:1403) CTTGGGCCCCTCTGGGGGCTGGGTT (FRAG. NO:1394)(SEQ. ID NO:1404) TCCTGCTGCGCCTGGGCGCTG (FRAG. NO:1395)(SEQ. ID NO:1405) GCGTCTTGGGGTGC (FRAG. NO:1396)(SEQ. ID NO:1406) GGGGCCGGGGGCCGGGGG (FRAG. NO:1397)(SEQ. ID NO:1407) GCCGCTGTTCGTGGGCCTGGG (FRAG. NO:1398)(SEQ. ID NO:1408) 10 GGTGCCTGTGGCTGCC (FRAG. NO:1399)(SEQ. ID NO:1409) GGTTGCCCCGGTTGGTGGC (FRAG. NO:1400)(SEQ. ID NO:1410) GCCGTCCTGCCGGT (FRAG. NO:1401)(SEQ. ID NO:1411) CGTTGGCTGGGTCCCCCCGC (FRAG. NO:1402)(SEQ. ID NO:1412) CCGTTTCCTGGGGTCC (FRAG. NO:1403)(SEQ. ID NO:1413) 15 GCGTGGGGTGCTCC (FRAG. NO:1404)(SEQ. ID NO:1414) GGTTCCTCGTGCCG (FRAG. NO:1405)(SEQ. ID NO:1415) CTGCTGCCTTGTCTTTCC (FRAG. NO:1406)(SEQ. ID NO:1416) GGCCGTGGCGGCGTGGTGCC (FRAG. NO:1407)(SEQ. ID NO:1417) GCCCCCCTGGCCTTCTGCTC (FRAG. NO:1408)(SEQ. ID NO:1418) 20 GGGGTCTGGCTGGT (FRAG. NO:1409)(SEQ. ID NO:1419) TGCCGGTGCCCTTGGCGGC (FRAG. NO:1410)(SEQ. 1D NO:1420) GGTCTTCTTCCTGGTG (FRAG. NO:1411)(SEQ. ID NO:1421) GCTCTGGGCCCGGCCGGTCTCGG (FRAG. NO:1412)(SEQ. ID NO:1422) GCGTCTCGTGTTCG (FRAG. NO:1413)(SEQ. ID NO:1423) 25 CTCTTGTGCTGTTCCGGCCG (FRAG. NO:1414)(SEQ. ID NO:1424) CTCCTTCCTCTCCGCCGCC (FRAG. NO:1415)(SEQ. ID NO:1425) GCCGCTCCCCGCCC (FRAG. NO:1416)(SEQ. ID NO:1426) GCTCGTCGCCCTGGCCC (FRAG. NO:1417)(SEQ. ID NO:1427) GGCCTCCTCCTGGCCGC (FRAG. NO:1418)(SEQ. ID NO:1428) 30 TGTCTCGGGCGGCGGCCTTGGC (FRAG. NO:1419)(SEQ. ID NO:1429) GCTCCGTTTGGGGCTG (FRAG. NO:1420)(SEQ. ID NO:1430) CCTCTGGCGCTTCC (FRAG. NO:1421)(SEQ. ID NO:1431) GGCCCTCGGCCTGGGCGCTC (FRAG. NO:1422)(SEQ. ID NO:1432) TCTTCCGCCTGTGC (FRAG. NO:1423)(SEQ. ID NO:1433) 35 TGGTGGCCCTCGTGG (FRAG. NO:1424)(SEQ. ID NO:1434) GCCCCTCCTGGCCTCCGGTGTCC (FRAG. NO:1425)(SEQ. ID NO:1435) TGTGGTCCCCGGCTGGT (FRAG. NO:1426)(SEQ. ID NO:1436) GGCCGGGCCGGTTGGGCGGGC (FRAG. NO:1427)(SEQ. ID NO:1437) GTGGGCGCCGGCGGTCCTCC (FRAG. NO:1428)(SEQ. ID NO:1438) 40 GGGCTGCCCTTCTCC (FRAG. NO:1429)(SEQ. ID NO:1439) GCCGGGGGTCCCGC (FRAG. NO:1430)(SEQ. ID NO:1440)
GCTCCTGCTGTTCCCTGGGCTCTTCTGCC (FRAG. NO:1431)(SEQ. ID NO:1441) TCTCTCCTGGGTGGGTGCCG (FRAG. NO:1432)(SEQ. ID NO:1442) GGGTCTCCGGGCTTG (FRAG. NO:1433)(SEQ. ID NO:1443) 45 CCCCGCGCTGCTGGGCGTTCTGC (FRAG. NO:1434)(SEQ. ID NO:1444) GGTCTTGGGGTTGTC (FRAG. NO:1435)(SEQ. ID NO:1445) TGTGGCCCCGCTCG (FRAG. NO:1436)(SEQ. ID NO:1446) TGTCGCCCTCCGTCGCC (FRAG. NO:1437)(SEQ. ID NO:1447) CGTCGCCGGCCTCGTCC (FRAG. NO:1438)(SEQ. ID NO:1448) 50 CCTCCTGGGTGCGC (FRAG. NO:1439)(SEQ. ID NO:1449) GGCGGGCTGGTCCT (FRAG. NO:1440)(SEQ. ID NO:1450) GGCGTTTTGCTCCTTCCTGG (FRAG. NO:1441)(SEQ. ID NO:1451) 5'-GCGTCTTGGGGTGCBGGCCCBTCCTGCTGCGCCTGGGCGCTG-3' (FRAG. NO:1896) (SEQ. ID NO: 1907)

55 Inducible Nitric Oxide Synthase Antisense Oligonucleotide Fragments

5'-CTGCCCCBGT TTTTGBTCCT CBCBTGCCGT GGGGBGGBCB BTGGCTGCCT CCCCGGGGTT TCTGCTGCTT GCTGCTTCTT TCCCGTCTCC CTTCTTTCCC GTCTCCTTTT TGCCTCTTTT GGTTCCTGTT GTTTCTGGCC TGCTTGTGG CGGCTTGTGC GTTTCCTCTC TCTTCTCTTG GGTCCCGCT TCTCGTCCTG CCTTTTCCTG TCTCTGTCGC GCCGTTCCTC CTCCGGCGTC CTCCTGCCCT GTGCTGTTTG CCTCGGGTGG TGCGGGTCCC GGTGCTCCC CGGCGGGCC GCTGGTTGC TGGGCCTGTC TGGTGGGGT TGGGGGGGT TCTTGTGGGC TGTGGTGGGC TCTTCTGTGG CCTTTGGGGC TGTGGTGGC CTGGCTCCTT GGGCCGTTCCCCGC CCCCCTTCTG GGCCGGTGGC CTGGCTCCTT GTGGGCGCTT CTGGCGCTT CTGGCCTCTT CTTCGCCTCG TGGCTGCTG GCTGC-3' (FRAG. NO:1897) (SEQ. ID NO: 1908)

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5'-CCCCGGGG-3' (FRAG. NO:1898) (SEQ. ID NO: 1909)
       5'-GGGGCCGCTGGG-3' (FRAG. NO:1899) (SEQ. ID NO:1910)
       5'-GGGGGTGTGG-3' (FRAG. NO:1900) (SEQ. ID NO: 1911)
       5'-CTGCCTCCCGGGGT-3' (FRAG. NO:1442)(SEQ. ID NO:1452)
       5'-TTCTGCTGCTTGCTG-3' (FRAG. NO:1443)(SEQ. ID NO:1453)
       5'-CTTCTTTCCCGTCTCC-3' (FRAG. NO:1444)(SEQ. ID NO:1454)
       5'-CTTCTTTCCCGTCTCC-3' (FRAG. NO:1445)(SEQ. ID NO:1455)
       5'-TTTTTGCCTCTTTG-3' (FRAG. NO:1446)(SEQ. ID NO:1456)
       5'-GGTTCCTGTTGTTTCT-3' (FRAG. NO:1447)(SEQ. ID NO:1457)
       5'-GGCCTGCTTGGTGGCG-3' (FRAG. NO:1448)(SEQ. ID NO:1458)
10
       5'-GCTTGTGCGTTTCC-3' (FRAG. NO:1449)(SEQ. ID NO:1459)
       5'-TCTCTCTCTCTCTGGGTCTCCGCCTTCTCGTCCTGCC-3' (FRAG. NO:1450)(SEQ. ID NO:1460)
       5'-TTTTCCTGTCTCTGTCGC-3' (FRAG. NO:1451)(SEQ. ID NO:1461)
       5'-GCCGTTCCTCCTCC-3' (FRAG, NO:1452)(SEQ. ID NO:1462)
       5'-GGCGTCCTCCTGCCC-3' (FRAG. NO:1453)(SEQ. ID NO:1463)
15
        5'-TGTGCTGTTTGCCTCGG-3' (FRAG. NO:1454)(SEQ. ID NO:1464)
        5'-GTGGTGCGGGTCCC-3' (FRAG. NO:1455)(SEQ. ID NO:1465)
        5'-GGTGCTCCCCGGC-3' (FRAG. NO:1456)(SEQ. ID NO:1466)
        5'-GGGCCGGCTGGTTGCCTGGGC-3' (FRAG. NO:1457)(SEQ. ID NO:1467)
        5'-CTGTCTGGTGGGGTGTGGGGCC-3' (FRAG. NO:1458)(SEQ. ID NO:1468)
20
        5'-GCTGGGTTGGGGGTGTGGTG-3' (FRAG. NO:1459)(SEQ. ID NO:1469)
        5'-GGCTCTTCTGTGGCC-3' (FRAG. NO:1460)(SEQ. ID NO:1470)
        5'-TGTGGGGCTGTTGGTG-3' (FRAG. NO:1461)(SEQ. ID NO:1471)
        5'-TCTCTGTGGGCGTGTG-3' (FRAG. NO:1462)(SEQ. ID NO:1472)
        5'-CTGGGTCTTGGGGCTTC-3' (FRAG. NO:1463)(SEQ. ID NO:1473)
25
        5'-CTCCCTTGTGCTGGG-3' (FRAG. NO:1464)(SEQ. ID NO:1474)
        5'-TGCGGCCTCCCCGC-3' (FRAG. NO:1465)(SEQ. ID NO:1475)
        5'-CCCCCTTCTGGGCC-3' (FRAG. NO:1466)(SEQ. ID NO:1476)
        5'-GGTGGCCTGGCTCCTTGTGG-3' (FRAG. NO:1467)(SEQ. ID NO:1477)
        5'-GCGCTTCTGGCTCTTG-3' (FRAG. NO:1468)(SEQ. ID NO:1478)
30
        5'-CCCTGTCCTTCTTCGCCTCGT-3' (FRAG. NO:1469)(SEQ. ID NO:1479)
        5'-GGCTGCTGGGCTGC-3' (FRAG. NO:1470)(SEQ. ID NO:1480)
        5'-CTGCCCCBGTTTTTGBTCCTCBCBTGCCGTGGGGBGGBCBBTGG-3' (FRAG. NO:1901) (SEQ. ID NO: 1912)
        NF-kB Antisense Oligonucleotide Fragments
        5'-CGGCCCTTCT CACTGGAGGC ACCGGGCAGT CCTCCATGGG AGGGTTGGGC TTGGCCGGGG CTGCCCGGTG CCTCCTCTTG
 35
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- GCTGGTCCCT CGTTGTCCTT GGGCCCCGC TCCCGCTGCT CGGCCTCCGT GTTCTTTGGC CTCTTGCTCC GCCTGCTGTC TIGTCCCGTC CCCTCCTCGC TTGCGTTTCC CTCTTCCTTG TCTTCCAGGC CTTCCTCCGC TTCCGCTGCT GGGGCCCGCG CCGGGGGGC GCTCGGCTCC GCGGCTTCCT CCCCGGCTGG GGGGTCCTGG TCTCCGGGGC CTGCGGCTCG CGGGCTCGGG GCTGCGTGCG CCGCGCGGG CGTCCGCGGT GGGTGGCGCT GTCCCGCCGT GGTGTGTCTC CGTTCTCGTC CTGCGCCGTC CTGGTCTGCC CGTGGGGTCC TGGGGGTGGT GGGGGGCGTC TGGTGCCTCG TCTGCCCCGT GGGGCTTCGG GCTCGGGGCT 40 GTTCGTCCCC CCTGCCGCTC TGTGGCCTCC GGGGCTCCTC GTTTTCGCTG CTTCGGGCTGT CCTTCTCGGC GTGTGGCCCC GGGTCCCGGC CCTGCTGGGC TGGGCGGGGT CGCTGCCCTG GGCTTCTGGC CCGTCTGGTT GTCTGTCGGT GCTTGTCTCG
- GGTTTCTGGC CTCTGTGCTG GGCGCTTCTC TGCCTCCTGC TCCGCCCTCC TGGTGGCTCG GCTGGGGGTG CCCGTGCGGG GGTGGGTGTG GGGTGTTTTC GGGGTCCTCC CCTTCCC-3' (FRAG. NO:1902) (SEQ. ID NO:1913)
- 5'-GGGCGGGGTCGC-3' (FRAG. NO:1903) (SEQ. ID NO:1914) 45
 - 5'-GCGCCGTCC-3' (FRAG. NO:1904) (SEQ. ID NO:1915)
 - 5'-GGGCGTGGTGG-3' (FRAG. NO:1905) (SEQ. ID NO:1916)
 - 5'-GTTGGGCTTGGCCGGGG-3' (FRAG. NO:1471)(SEQ. ID NO:1481)
 - 5'-CTGCCCGGTGCCTCC-3' (FRAG. NO:1472)(SEQ. ID NO:1482)
- 5'-TCTTGGCTGGTCCCTCGT-3' (FRAG. NO:1473)(SEQ. ID NO:1483) 50
 - 5'-TGTCCTTGGGCCCC-3' (FRAG. NO:1474)(SEQ. ID NO:1484)
 - 5'-GCTCCCGCTGCTCGGCCTCCGT-3' (FRAG. NO:1475)(SEQ. ID NO:1485)
 - 5'-GTTCTTTGGCCTCTTGCTCC-3' (FRAG. NO:1476)(SEQ. ID NO:1486)
 - 5'-GCCTGCTGTCTTGTCC-3' (FRAG. NO:1477)(SEQ. ID NO:1487)
- 5'-CGTCCCCTCCTCGCTTGCGTTTC-3' (FRAG. NO:1478)(SEQ. ID NO:1488) 55 5'-CCTCTTCCTTGTCTTCCA-3' (FRAG. NO:1479)(SEQ. ID NO:1489)
 - 5'-GGCCTTCCTCCGCTTCCGCTGC-3' (FRAG. NO:1480)(SEQ. ID NO:1490)
 - 5'-TGGGGCCCGCCGC-3' (FRAG. NO:1481)(SEQ. ID NO:1491)
- 5'-GGGGGCGCTCCGCGGCTTCCTCCCCGG-3' (FRAG. NO:1482)(SEQ. ID NO:1492) 5'-CTGGGGGGTCCTGG-3' (FRAG. NO:1483)(SEQ. ID NO:1493) 60
 - 5'-TCTCCGGGGCCTGCGGCTCGC-3' (FRAG. NO:1484)(SEQ. ID NO:1494)
 - 5'-GGGCTCGGGGCTGCGTGCGCC-3' (FRAG. NO:1485)(SEQ. ID NO:1495)

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5'-GCGCGCGCGCCTCCGCGGTG-3' (FRAG. NO:1486)(SEQ. ID NO:1496)
       5'-GGTGGCGCTGTCCCGCC-3' (FRAG. NO:1487)(SEQ. ID NO:1497)
       5'-GTGGTGTGTCTCCGTTCTCGTCCTGCGCCGTC-3' (FRAG. NO:1488)(SEQ. ID NO:1498)
       5'-CTGGTCTGCCCGTGG-3' (FRAG. NO:1489)(SEQ. ID NO:1499)
       5'-GGTCCTGGGCGTGGTGG-3' (FRAG. NO:1490)(SEQ. ID NO:1500)
       5'-GGGGCGTCTGGTGC-3' (FRAG. NO:1491)(SEQ. ID NO:1501)
       5'-CTCGTCTGCCCCGTG-3' (FRAG. NO:1492)(SEQ. ID NO:1502)
       5'-GGGCTTCGGGCTCGG-3' (FRAG. NO:1493)(SEQ. ID NO:1503)
       5'-GGCTGTTCGTCCCCCTGCCGCTCTGTGGCCTCC-3' (FRAG. NO:1494)(SEQ. ID NO:1504)
       5'-GGGGCTCCTCGTTTTC-3' (FRAG. NO:1495)(SEQ. ID NO:1505)
10
       5'-GCTGCTTCGGGTGTCCTTCTC-3' (FRAG. NO:1496)(SEQ. ID NO:1506)
       5'-GGCGTGTGGCCCCGG-3' (FRAG. NO:1497)(SEQ. ID NO:1507)
       5'-GTCCCGGCCCTGCTGGGCTGGGCGGGGTC-3' (FRAG, NO:1498)(SEQ, ID NO:1508)
       5'-GCTGCCCTGGGCTTCTGGCCCGTCT-3' (FRAG. NO:1499)(SEQ. ID NO:1509)
       5'-GGTTGTCTGTCGGT-3' (FRAG. NO:1500)(SEQ. ID NO:1510)
15
       5'-GCTTGTCTCGGGTTTCTGG-3' (FRAG. NO:1501)(SEQ. ID NO:1511)
       5'-CCTCTGTGCTGGGC-3' (FRAG. NO:1502)(SEQ. ID NO:1512)
       5'-GCTTCTCTGCCTCCTGCTCC-3' (FRAG. NO:1503)(SEQ. ID NO:1513)
       5'-GCCCTCCTGGTGGCTC-3' (FRAG. NO:1504)(SEQ. ID NO:1514)
       5'-GGCTGGGGGTGCCCGTGCG-3' (FRAG. NO:1505)(SEQ. ID NO:1515)
20
       5'-GGGGTGGGTGTGGGGTGTT-3' (FRAG. NO:1506)(SEQ. ID NO:1516)
       5'-TTCGGGGTCCTCCCCTTCCC-3' (FRAG. NO:1507)(SEQ. ID NO:1517)
       5'-CGGCCCTTCTCACTGGAGGCACCGGGCAGTCCTCCATGGGAGG-3' (FRAG. NO:1906) (SEQ. ID NO:1917)
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Human Major Basic Protein Anti-sense Oligonucleotide Fragments

- 30 5'-GGG GGA GTT-3' (FRAG. ID:1908) (SEQ. ID NO:1919)
 5'-G CCC TGG GCC C-3' (FRAG. ID:1909) (SEQ. ID NO:1920)
 5'-GTT TCA TCT TGG CTT TAT CC-3' (FRAG. NO:1508) (SEQ. ID NO:1518)
 5'-TCT CCC CTT GTT CCT CCC C-3' (FRAG. NO:1509)(SEQ. ID NO:1519)

5'-TCT CCT GCT CTG GRG TCT CCT C-3' (FRAG. NO:1510)(SEQ. ID NO:1520)

5'-TTC CCT CCC TCC CCT GCC-3' (FRAG. NO:1511)(SEQ. ID NO:1521)

5'-GTG TTG TCT GTG GGT GTC C-3' (FRAG. NO:1512)(SEQ. ID NO:1522)

5'-GTT TCG CTC TTG TTG CCC-3' (FRAG. NO:1513)(SEQ. ID NO:1523)

5' TGG GCC CTT CCC TGC TGG-3' (FRAG. NO:1514)(SEO. ID NO:1524)

5'-TGG GCC CTT CCC TGC TGG-3' (FRAG. NO:1514)(SEQ. ID NO:1524) 5'-GGG GGA GTT TCA TCT TGG-3' (FRAG. NO:1515)(SEQ. ID NO:1525)

Human Eosinophil Major Basic Protein Fragments

- 5'-GGG GGB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1516)(SEQ. ID NO:1526) 5'-GGG GGB GTT TCB TCT TGG CTT-3' (FRAG. NO:1517)(SEQ. ID NO:1527) 5'-GGG GGB GTT TCB TCT TGG CT-3' (FRAG. NO:1518)(SEQ. ID NO:1528)
- 5'-GGG GGB GTT TCB TCT TGG C-3' (FRAG. NO:1519)(SEQ. ID NO:1529) 5'-GGG GGB GTT TCB TCT TGG-3' (FRAG. NO:1520)(SEQ. ID NO:1530)
 - 5'-GGG GGB GTT TCB TCT TG-3' (FRAG. NO:1521)(SEQ. ID NO:1531) 5'-GGG GGB GTT TCB TCT T-3' (FRAG. NO:1522)(SEQ. ID NO:1532)
- 5'-GGG GGB GTT TCB TCT-3' (FRAG. NO:1523)(SEQ. ID NO:1533)
 55 5'-GGG GGB GTT TCB TC-3' (FRAG. NO:1524)(SEQ. ID NO:1534)
 - 5'-GGG GGB GTT TCB T-3' (FRAG. NO:1525)(SEQ. ID NO:1535)
 - 5'-GGG GGB GTT TCB-3' (FRAG. NO:1526)(SEQ. ID NO:1536) 5'-GG GGB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1527)(SEQ. ID NO:1537)
 - 5'-GG GGB GTT TCB TCT TGG CTT-3' (FRAG. NO:1528)(SEQ. ID NO:1538)
- 60 5'-GG GGB GTT TCB TCT TGG CT-3' (FRAG. NO:1529)(SEQ. ID NO:1539)

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5'-GG GGB GTT TCB TCT TGG C-3' (FRAG. NO:1530)(SEQ. ID NO:1540)
       5'-GG GGB GTT TCB TCT TGG-3' (FRAG. NO:1531)(SEQ. ID NO:1541)
       5'-GG GGB GTT TCB TCT TG-3' (FRAG. NO:1532)(SEQ. ID NO:1542)
       5'-GG GGB GTT TCB TCT T-3' (FRAG. NO:1533)(SEQ. ID NO:1543)
       5'-GG GGB GTT TCB TCT-3' (FRAG. NO:1534)(SEQ. ID NO:1544)
        5'-GG GGB GTT TCB TC-3' (FRAG. NO:1535)(SEQ. ID NO:1545)
        5'-GG GGB GTT TCB T-3' (FRAG. NO:1536)(SEQ. ID NO:1546)
        5'-G GGB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1537)(SEQ. ID NO:1547)
        5'-G GGB GTT TCB TCT TGG CTT-3' (FRAG. NO:1538)(SEQ. ID NO:1548)
        5'-G GGB GTT TCB TCT TGG CT-3' (FRAG. NO:1539)(SEQ. ID NO:1549)
10
        5'-G GGB GTT TCB TCT TGG C-3' (FRAG. NO:1540)(SEQ. ID NO:1550)
        5'-G GGB GTT TCB TCT TGG-3' (FRAG. NO:1541)(SEQ. ID NO:1551)
        5'-G GGB GTT TCB TCT TG-3' (FRAG. NO:1542)(SEQ. ID NO:1552)
        5'-G GGB GTT TCB TCT T-3' (FRAG. NO:1543)(SEQ. ID NO:1553)
        5'-G GGB GTT TCB TCT-3' (FRAG. NO:1544)(SEQ. ID NO:1554)
15
        5'-G GGB GTT TCB TC-3' (FRAG. NO:1545)(SEQ. ID NO:1555)
        5'-GGB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1546)(SEQ. ID NO:1556)
        5'-GGB GTT TCB TCT TGG CTT-3' (FRAG. NO:1547)(SEQ. ID NO:1557)
        5'-GGB GTT TCB TCT TGG CT-3' (FRAG. NO:1548)(SEQ. ID NO:1558)
        5'-GGB GTT TCB TCT TGG C-3' (FRAG. NO:1549)(SEQ. ID NO:1559)
20
        5'-GGB GTT TCB TCT TGG-3' (FRAG. NO:1550)(SEQ. ID NO:1560)
        5'-GGB GTT TCB TCT TG-3' (FRAG. NO:1551)(SEQ. ID NO:1561)
        5'-GGB GTT TCB TCT T-3' (FRAG. NO:1552)(SEQ. ID NO:1562)
        5'-GGB GTT TCB TCT-3' (FRAG. NO:1553)(SEQ. ID NO:1563)
        5'-GB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1554)(SEQ. ID NO:1564)
25
        5'-GB GTT TCB TCT TGG CTT-3' (FRAG. NO:1555)(SEQ. ID NO:1565)
         5'-GB GTT TCB TCT TGG CT-3' (FRAG. NO:1556)(SEQ. ID NO:1566)
         5'-GB GTT TCB TCT TGG C-3' (FRAG. NO:1557)(SEQ. ID NO:1567)
         5'-GB GTT TCB TCT TGG-3' (FRAG. NO:1558)(SEQ. ID NO:1568)
         5'-GB GTT TCB TCT TG-3' (FRAG. NO:1559)(SEQ. ID NO:1569)
30
         5'-GB GTT TCB TCT T-3' (FRAG. NO:1560)(SEQ. ID NO:1570)
         5'-B GTT TCB TCT TGG CTT T-3' (FRAG. NO:1561)(SEQ. ID NO:1571)
         5'-B GTT TCB TCT TGG CTT-3' (FRAG. NO:1562)(SEQ. ID NO:1572)
5'-B GTT TCB TCT TGG CTT-3' (FRAG. NO:1563)(SEQ. ID NO:1573)
         5'-B GTT TCB TCT TGG CT-3' (FRAG. NO:1564)(SEQ. ID NO:1574)
 35
         5'-B GTT TCB TCT TGG C-3' (FRAG. NO:1565)(SEQ. ID NO:1575)
         5'-B GTT TCB TCT TGG-3' (FRAG. NO:1565)(SEQ. ID NO:1576)
         5'-B GTT TCB TCT TG-3' (FRAG. NO:1567)(SEQ. ID NO:1577)
         5'-GTT TCB TCT TGG CTT T-3' (FRAG. NO:1568)(SEQ. ID NO:1578)
         5'-GTT TCB TCT TGG CTT-3' (FRAG. NO:1569)(SEQ. ID NO:1579)
 40
         5'-GTT TCB TCT TGG CT-3' (FRAG. NO:1570)(SEQ. ID NO:1580)
         5'-GTT TCB TCT TGG C-3' (FRAG. NO:1571)(SEQ. ID NO:1581)
         5'-GTT TCB TCT TGG-3' (FRAG. NO:1572)(SEQ. ID NO:1582)
         5'-TT TCB TCT TGG CTT T-3' (FRAG. NO:1573)(SEQ. ID NO:1583)
         5'-TT TCB TCT TGG CTT-3' (FRAG. NO:1574)(SEQ. ID NO:1584)
 45
         5'-TT TCB TCT TGG CT-3' (FRAG. NO:1575)(SEQ. ID NO:1585)
         5'-TT TCB TCT TGG C-3' (FRAG. NO:1576)(SEQ. ID NO:1586)
         5'-T TCB TCT TGG CTT T-3' (FRAG. NO:1577)(SEQ. ID NO:1587)
         5'-T TCB TCT TGG CTT-3' (FRAG. NO:1578)(SEQ. ID NO:1588)
         5'-T TCB TCT TGG CT-3' (FRAG. NO:1579)(SEQ. ID NO:1589)
 50
          5'-TCB TCT TGG CTT T-3' (FRAG. NO:1580)(SEQ. ID NO:1590)
          5'-TCB TCT TGG CTT-3' (FRAG. NO:1581)(SEQ. ID NO:1591)
          5'-GGG GGB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1582)(SEQ. ID NO:1592)
          5'-GG GGB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1583)(SEQ. ID NO:1593)
          5'-G GGB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1584)(SEQ. ID NO:1594)
 55
          5'-GGB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1585)(SEQ. ID NO:1595)
          5'-GB GTT TCB TCT TGG CTT T-3' (FRAG. NO:1586)(SEQ. ID NO:1596)
          5'-B GTT TCB TCT TGG CTT T-3' (FRAG. NO:1587)(SEQ. ID NO:1597)
          5'-GTT TCB TCT TGG CTT T-3' (FRAG. NO:1588)(SEQ. ID NO:1598)
          5'-TT TCB TCT TGG CTT T-3' (FRAG. NO:1589)(SEQ. ID NO:1599)
  60
          5'-T TCB TCT TGG CTT T-3' (FRAG. NO:1590)(SEQ. ID NO:1600)
          5'-TCB TCT TGG CTT T-3' (FRAG. NO:1591)(SEQ. ID NO:1601)
          5'-CB TCT TGG CTT T-3' (FRAG. NO:1592)(SEQ. ID NO:1602)
          5'-GGG GGB GTT TCB TCT TGG CTT-3' (FRAG. NO:1593)(SEQ. ID NO:1603)
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5'-GG GGB GTT TCB TCT TGG CTT-3' (FRAG. NO:1594)(SEQ. ID NO:1604)
       5'-G GGB GTT TCB TCT TGG CTT-3' (FRAG. NO:1595)(SEQ. ID NO:1605)
       5'-GGB GTT TCB TCT TGG CTT-3' (FRAG. NO:1596)(SEQ. ID NO:1606)
       5'-GB GTT TCB TCT TGG CTT-3' (FRAG. NO:1597)(SEQ. ID NO:1607)
       5'-B GTT TCB TCT TGG CTT-3' (FRAG. NO:1598)(SEQ. ID NO:1608)
       5'-GTT TCB TCT TGG CTT-3' (FRAG. NO:1599)(SEQ. ID NO:1609)
       5'-TT TCB TCT TGG CTT-3' (FRAG. NO:1600)(SEQ. ID NO:1610)
       5'-T TCB TCT TGG CTT-3' (FRAG. NO:1601)(SEQ. ID NO:1611)
       5'-TCB TCT TGG CTT-3' (FRAG. NO:1602)(SEQ. ID NO:1612)
       5'-GGG GGB GTT TCB TCT TGG CT-3' (FRAG. NO:1603)(SEQ. ID NO:1613)
10
       5'-GG GGB GTT TCB TCT TGG CT-3' (FRAG. NO:1604)(SEQ. ID NO:1614)
       5'-G GGB GTT TCB TCT TGG CT-3' (FRAG. NO:1605)(SEQ. ID NO:1615)
       5'-GGB GTT TCB TCT TGG CT-3' (FRAG. NO:1606)(SEQ. ID NO:1616)
       5'-GB GTT TCB TCT TGG CT-3' (FRAG. NO:1607)(SEQ. ID NO:1617)
        5'-B GTT TCB TCT TGG CT-3' (FRAG. NO:1608)(SEQ. ID NO:1618)
15
        5'-GTT TCB TCT TGG CT-3' (FRAG. NO:1609)(SEQ. ID NO:1619)
        5'-TT TCB TCT TGG CT-3' (FRAG. NO:1610)(SEQ. ID NO:1620)
        5'-T TCB TCT TGG CT-3' (FRAG. NO:1611)(SEQ. ID NO:1621)
        5'-GGG GGB GTT TCB TCT TGG C-3' (FRAG. NO:1612)(SEQ. ID NO:1622)
        5'-GG GGB GTT TCB TCT TGG C-3' (FRAG. NO:1613)(SEQ. ID NO:1623)
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        5'-G GGB GTT TCB TCT TGG C-3' (FRAG. NO:1614)(SEQ. ID NO:1624)
        5'-GGB GTT TCB TCT TGG C-3' (FRAG. NO:1615)(SEQ. ID NO:1625)
        5'-GB GTT TCB TCT TGG C-3' (FRAG. NO:1616)(SEQ. ID NO:1626)
        5'-B GTT TCB TCT TGG C-3' (FRAG. NO:1617)(SEQ. ID NO:1627)
        5'-GTT TCB TCT TGG C-3' (FRAG. NO:1618)(SEQ. ID NO:1628)
25
        5'-TT TCB TCT TGG C-3' (FRAG. NO:1619)(SEQ. ID NO:1629)
        5'-GGG GGB GTT TCB TCT TGG-3' (FRAG. NO:1620)(SEQ. ID NO:1630)
        5'-GG GGB GTT TCB TCT TGG-3' (FRAG. NO:1621)(SEQ. ID NO:1631)
        5'-G GGB GTT TCB TCT TGG-3' (FRAG. NO:1622)(SEQ. ID NO:1632)
        5'-GGB GTT TCB TCT TGG-3' (FRAG. NO:1623)(SEQ. ID NO:1633)
30
         5'-GB GTT TCB TCT TGG-3' (FRAG. NO:1624)(SEQ. ID NO:1634)
         5'-B GTT TCB TCT TGG-3' (FRAG. NO:1625)(SEQ. ID NO:1635)
         5'-GTT TCB TCT TGG-3' (FRAG. NO:1626)(SEQ. ID NO:1636)
         5'-GGG GGB GTT TCB TCT TG-3' (FRAG. NO:1627)(SEQ. ID NO:1637)
         5'-GG GGB GTT TCB TCT TG-3' (FRAG. NO:1628)(SEQ. ID NO:1638)
 35
         5'-G GGB GTT TCB TCT TG-3' (FRAG. NO:1629)(SEQ. ID NO:1639)
         5'-GGB GTT TCB TCT TG-3' (FRAG. NO:1630)(SEQ. ID NO:1640)
         5'-GB GTT TCB TCT TG-3' (FRAG. NO:1631)(SEQ. ID NO:1641)
         5'-B GTT TCB TCT TG-3' (FRAG. NO:1632)(SEQ. ID NO:1642)
         5'-GGG GGB GTT TCB TCT T-3' (FRAG. NO:1633)(SEQ. ID NO:1643)
 40
         5'-GG GGB GTT TCB TCT T-3' (FRAG. NO:1634)(SEQ. ID NO:1644)
         5'-G GGB GTT TCB TCT T-3' (FRAG. NO:1635)(SEQ. ID NO:1645)
5'-G GGB GTT TCB TCT T-3' (FRAG. NO:1636)(SEQ. ID NO:1646)
         5'-GGB GTT TCB TCT T-3' (FRAG. NO:1637)(SEQ. ID NO:1647)
         5'-GB GTT TCB TCT T-3' (FRAG. NO:1638)(SEQ. ID NO:1648)
 45
         5'-GGG GGB GTT TCB TCT-3' (FRAG. NO:1639)(SEQ. ID NO:1649)
         5'-GG GGB GTT TCB TCT-3' (FRAG. NO:1640)(SEQ. 1D NO:1650)
         5'-G GGB GTT TCB TCT-3' (FRAG. NO:1641)(SEQ. ID NO:1651)
         5'-GGB GTT TCB TCT-3' (FRAG. NO:1642)(SEQ. ID NO:1652)
          5'-GGG GGB GTT TCB TC-3' (FRAG. NO:1643)(SEQ. ID NO:1653)
 50
          5'-GG GGB GTT TCB TC-3' (FRAG. NO:1644)(SEQ. ID NO:1654)
          5'-G GGB GTT TCB TC-3' (FRAG. NO:1645)(SEQ. ID NO:1655)
          5'-GGG GGB GTT TCB T-3' (FRAG. NO:1646)(SEQ. ID NO:1656)
          5'-GG GGB GTT TCB T-3' (FRAG. NO:1647)(SEQ. ID NO:1657)
          5'-GGG GGB GTT TCB-3' (FRAG. NO:1648)(SEQ. ID NO:1658)
  55
          5'-TCT CCC CTT GTT CCT CCC C-3' (FRAG. NO:1649)(SEQ. ID NO:1659)
          5'-TCT CCT GCT CTG GTG TCT CCT C-3' (FRAG. NO:1650)(SEQ. ID NO:1660)
          5'-TTC CCT CCC TCC CCT GCC-3' (FRAG. NO:1651)(SEQ. ID NO:1661)
          5'-GTG TTG TCT GTG GGT GTC C-3' (FRAG. NO:1652)(SEQ. ID NO:1662)
          5'-GTT TCG CTC TTG TTG CCC-3' -3' (FRAG. NO:1653)(SEQ. ID NO:1663)
  60
          5'-TGG GCC CTT CCC TGC TGG-3' (FRAG. NO:1654)(SEQ. ID NO:1664)
          5'-GGG GGB G-3' (FRAG. NO:1912)(SEQ. ID NO:1923)
          5'-GTG GGT GTC C-3' (FRAG. NO:1913) (SEQ. ID NO:1924)
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BP-1 Antisense Oligonucleotide Fragments

5'-CCGTGTTGTC BGTGGTGCTG CCCGTTTGBG GTBTGGCGCT CCBCCBBTTC CCTTTTCTCC TIGTTTTCCG TTTCTCTTGC CGTCTGTGGT T-3' (FRAG. NO:1914) (SEQ. ID NO:1925) 5'-CCCGTTTGBGGTBTGGC-3'(FRAG. NO:1915) (SEQ. ID NO:1926) 5'-GCTCCBCCBBTTCCCTTTTCTCC-3'(FRAG. NO:1916) (SEQ. ID NO:1927)

5'-TTGTTTTCCGTTTCTCTTG-3'(FRAG, NO:1917) (SEQ. ID NO:1928)

5'-CCGTCTGTGGTT-3'(FRAG. NO:1918) (SEQ. ID NO:1929)

5'-CCCGTTTGAGGTATGGC-3'(FRAG. NO:1919) (SEQ. ID NO:1930)

5'-GCTCCBCCAATTCCCTTTTCTCC-3'(FRAG. NO:1920) (SEQ. ID NO:1931)

C/EBPB Antisense Oligonucleotide Fragments 5'-GGGCCCBGCCCGCCCTTTTCTBGCCCC GGCC-3' (FRAG. NO:1921) (SEQ. ID NO:1932)

5'-GGGCCCBGCCCGCCGCCTTTTCTBGCCCC GGC-3' (FRAG. NO:1922) (SEQ. ID NO:1933)

5'-GGGCCCB GCCCGCCGCCTTTTCTBGCCCCGG-3' (FRAG. NO:1923) (SEQ. ID NO:1934)

5'-GGGCCCBGCCCGCCGCCTTTTCTBGCCCCG-3' (FRAG. NO:1924) (SEQ. ID NO:1935)

5'-GGGCCCBGCCCGCCCTTTTCTBGCCCC-3' (FRAG. NO:1925) (SEQ. ID NO:1936) 15

5'-GGGCCCBGCCCGCCGCCTTTTCTBGCCC-3' (FRAG, NO:1926) (SEQ. ID NO:1937)

5'-GGGCCCBGCCCGCCCTTTTCTBGCC-3' (FRAG. NO:1927) (SEQ. ID NO:1938)

5'-GGGCCCBGCCCGCCGCCTTTTCTBGC-3' (FRAG. NO:1928) (SEQ. ID NO:1939)

5'-GGGCCCBGCCCGCCGCCTTTTCTBG-3' (FRAG. NO:1929) (SEQ. ID NO:1940)

5'-GGGCCCBGCCCGCCGCCTTTTCTB-3' (FRAG. NO:1930) (SEQ. ID NO:1941)

5'-GGGCCCBGCCCGCCCTTTTCT-3' (FRAG. NO:1931) (SEQ. ID NO:1942)

5'-GGGCCCBGCCCGCCGCCTTTTC-3' (FRAG. NO:1932) (SEQ. ID NO:1943)

5'-GGGCCCBGCCCGCCGCCTTTT-3' (FRAG. NO:1933) (SEQ. ID NO:1944)

5'-GGGCCCBGCCCGCCCTTT-3' (FRAG. NO:1934) (SEQ. ID NO:1945)

5'-GGGCCCBGCCCGCCGCCTT-3' (FRAG. NO:1935) (SEQ. ID NO:1946)

5'-GGGCCCBGCCCGCCCT-3' (FRAG. NO:1936) (SEQ. ID NO:1947)

5'-GGGCCCBGCCCGCCGCC-3' (FRAG. NO:1937) (SEQ. ID NO:1948)

5'-GGGCCCBGCCCGCCGC-3' (FRAG. NO:1938) (SEQ. ID NO:1949)

5'-GGGCCCBGCCCGCCG-3' (FRAG. NO:1939) (SEQ. ID NO:1950)

5'-GGGCCCBGCCCGCC-3' (FRAG. NO:1940) (SEQ. ID NO:1951)

5'-GGGCCCBGCCCCGC-3' (FRAG. NO:1941) (SEQ. ID NO:1952)

5'-GGGCCCBGCCCCG-3' (FRAG. NO:1942) (SEQ. ID NO:1953) 5'-GGGCCCBGCCCC-3' (FRAG. NO:1943) (SEQ. ID NO:1954)

5'-GGGCCCBGCCC-3' (FRAG. NO:1944) (SEQ. ID NO:1955)

5'-GGCCCBGCCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1945) (SEQ. ID NO:1956) 35

5'-GCCCBGCCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1946) (SEQ. ID NO:1957)

5'-CCCBGCCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1947) (SEQ. ID NO:1958)

5'-CCBGCCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1948) (SEQ. ID NO:1959)

5'-CBGCCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1948) (SEQ. ID NO:1960)

5'-BGCCCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1950) (SEQ. ID NO:1961)

5'-GCCCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1951) (SEQ. ID NO:1962) 5'-CCCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1952) (SEQ. ID NO:1963)

5'-CCCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1953) (SEQ. ID NO:1964)

5'-CCGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1954) (SEQ. ID NO:1965)

5'-CGCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1955) (SEQ. ID NO:1966)

5'-GCCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1956) (SEQ. ID NO:1967)

5'-CCGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1957) (SEQ. ID NO:1968)

5'-CGCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1958) (SEQ. ID NO:1969)

5'-GCCTTTTCTBGCCCCGGC-3' (FRAG. NO:1959) (SEQ. ID NO:1970)

5'-CCTTTTCTBGCCCCGGC-3' (FRAG. NO:1960) (SEQ. ID NO:1971) 50

5'-CTTTTCTBGCCCCGGC-3' (FRAG. NO:1961) (SEQ. ID NO:1972)

5'-TTTTCTBGCCCCGGC-3' (FRAG. NO:1962) (SEQ. ID NO:1973)

5'-TTTCTBGCCCCGGC-3' (FRAG. NO:1963) (SEQ. ID NO:1974) 5'-TTCTBGCCCCGGC-3' (FRAG. NO:1964) (SEQ. ID NO:1975)

5'-TCTBGCCCCGGC-3' (FRAG. NO:1965) (SEQ. ID NO:1976) 55

5'-CTBGCCCCGGC-3' (FRAG. NO:1966) (SEQ. ID NO:1977)

5'-GCGBGGCTGTCBCCTCGCTGGGCCC-3' (FRAG. NO:1967) (SEQ. ID NO:1978)

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5'-GCGBGGCTGTCBCCTCGCTGGGCC-3' (FRAG. NO:1968) (SEQ. ID NO:1979)
       5'-GCGBGGCTGTCBCCTCGCTGGGC-3' (FRAG. NO:1969) (SEQ. ID NO:1980)
       5'-GCGBGGCTGTCBCCTCGCTGGG-3' (FRAG. NO:1970) (SEQ. ID NO:1981)
       5'-GCGBGGCTGTCBCCTCGCTGG-3' (FRAG. NO:1971) (SEQ. ID NO:1982)
       5'-GCGBGGCTGTCBCCTCGCTG-3' (FRAG. NO:1972) (SEQ. ID NO:1983)
5
       5'-GCGBGGCTGTCBCCTCGCT-3' (FRAG. NO:1973) (SEQ. ID NO:1984)
       5'-GCGBGGCTGTČBCCTCGC-3' (FRAG. NO:1974) (SEQ. ID NO:1985)
       5'-GCGBGGCTGTCBCCTCG-3' (FRAG. NO:1975) (SEQ. ID NO:1986)
       5'-GCGBGGCTGTCBCCTC-3' (FRAG. NO:1976) (SEQ. ID NO:1987)
       5'-GCGBGGCTGTCBCCT-3' (FRAG. NO:1977) (SEQ. ID NO:1988)
10
       5'-GCGBGGCTGTCBCC-3' (FRAG. NO:1978) (SEQ. ID NO:1989)
       5'-GCGBGGCTGTCBC-3' (FRAG. NO:1979) (SEQ. ID NO:1990)
       5'-GCGBGGCTGTCB-3' (FRAG. NO:1980) (SEQ. ID NO:1991)
       5'-GCGBGGCTGTC-3' (FRAG. NO:1981) (SEQ. ID NO:1992)
       5'-GCGBGGCTGT-3' (FRAG. NO:1982) (SEQ. ID NO:1993)
15
       5'-CGBGGCTGTCBCCTCGCTGGGCCC-3' (FRAG. NO:1983) (SEQ. ID NO:1994)
        5'-GBGGCTGTCBCCTCGCTGGGCCC-3' (FRAG. NO:1984) (SEQ. ID NO:1995)
        5'-BGGCTGTCBCCTCGCTGGGCCC-3' (FRAG. NO:1985) (SEQ. ID NO:1996)
        5'-GGCTGTCBCCTCGCTGGGCCC-3' (FRAG. NO:1986) (SEQ. ID NO:1997)
        5'-GCTGTCBCCTCGCTGGGCCC-3' (FRAG. NO:1987) (SEQ. ID NO:1998)
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        5'-CTGTCBCCTCGCTGGGCCC-3' (FRAG. NO:1988) (SEQ. ID NO:1999)
        5'-TGTCBCCTCGCTGGGCCC-3' (FRAG. NO:1989) (SEQ. ID NO:2000)
        5'-GTCBCCTCGCTGGGCCC-3' (FRAG. NO:1990) (SEQ. ID NO:2001)
        5'-TCBCCTCGCTGGGCCC-3' (FRAG. NO:1991) (SEQ. ID NO:2002)
        5'-CBCCTCGCTGGGCCC-3' (FRAG. NO:1992) (SEQ. ID NO:2003)
25
        5'-BCCTCGCTGGGCCC-3' (FRAG. NO:1993) (SEQ. ID NO:2004)
        5'-CCTCGCTGGGCCC-3' (FRAG. NO:1994) (SEQ. ID NO:2005)
        5'-CTCGCTGGGCCC-3' (FRAG. NO:1995) (SEQ. ID NO:2006)
        5'-TCGCTGGGCCC-3' (FRAG. NO:1996) (SEQ. ID NO:2007)
        5'-CGCTGGGCCC-3' (FRAG. NO:1997) (SEQ. ID NO:2008)
30
        5'-GCGCGGCCGTCBTGGCGGCCGTCGGGCCGGGC-3' (FRAG. NO:1998) (SEQ. ID NO:2009)
        5'-GCGCGGCCGTCBTGGCGGCGTCGGGCCGGG-3' (FRAG. NO:1999) (SEQ. ID NO:2010)
        5'-GCGCGGCCGTCBTGGCGGCGTCGGGCCGG-3' (FRAG. NO:2000) (SEQ. ID NO:2011)
        5'-GCGCGGCCGTCBTGGCGGCGTCGGGCCG-3' (FRAG. NO:2001) (SEQ. ID NO:2012)
        5'-GCGCGGCCGTCBTGGCGGCGTCGGGCC-3' (FRAG. NO:2002) (SEQ. ID NO:2013)
35
        5'-GCGCGGCCGTCBTGGCGGCGTCGGGC-3' (FRAG. NO:2003) (SEQ. 1D NO:2014)
        5'-GCGCGGCCGTCBTGGCGGCGTCGGG-3' (FRAG. NO:2004) (SEQ. ID NO:2015)
         5'-GCGCGGCCGTCBTGGCGGCGTCGG-3' (FRAG. NO:2005) (SEQ. ID NO:2016)
         5'-GCGCGGCCGTCBTGGCGGCGTCG-3' (FRAG. NO:2006) (SEQ. ID NO:2017)
         5'-GCGCGGCCGTCBTGGCGGCGTC-3' (FRAG. NO:2007) (SEQ. ID NO:2018)
 40
         5'-GCGCGGCCGTCBTGGCGGCGT-3' (FRAG. NO:2008) (SEQ. ID NO:2019)
         5'-GCGCGGCCGTCBTGGCGGCG-3' (FRAG. NO:2009) (SEQ. ID NO:2020)
         5'-GCGCGGCCGTCBTGGCGGC-3' (FRAG. NO:2010) (SEQ. ID NO:2021)
         5'-GCGCGGCCGTCBTGGCGG-3' (FRAG. NO:2011) (SEQ. ID NO:2022)
         5'-GCGCGGCCGTCBTGGCG-3' (FRAG. NO:2012) (SEQ. ID NO:2023)
 45
         5'-GCGCGGCCGTCBTGGC-3' (FRAG. NO:2013) (SEQ. ID NO:2024)
         5'-GCGCGGCCGTCBTGG-3' (FRAG. NO:2014) (SEQ. ID NO:2025)
         5'-GCGCGGCCGTCBTG-3' (FRAG. NO:2015) (SEQ. ID NO:2026)
         5'-GCGCGGCCGTCBT-3' (FRAG. NO:2016) (SEQ. ID NO:2027)
         5'-GCGCGGCCGTCB-3' (FRAG. NO:2017) (SEQ. ID NO:2028)
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         5'-GCGCGGCCGTC-3' (FRAG. NO:2018) (SEQ. ID NO:2029)
         5'-GCGCGGCCGT-3' (FRAG. NO:2019) (SEQ. ID NO:2030)
         5'-CGCGGCCGTCBTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2020) (SEQ. ID NO:2031)
          5'-GCGGCCGTCBTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2021) (SEQ. ID NO:2032)
          5'-CGGCCGTCBTGGCGGCCGTCGGGCCGGGC-3' (FRAG. NO:2022) (SEQ. ID NO:2033)
  55
          5'-GGCCGTCBTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2023) (SEQ. ID NO:2034)
          5'-GCCGTCBTGGCGGCCGTCGGGCCGGCC-3' (FRAG. NO:2024) (SEQ. ID NO:2035)
          5'-CCGTCBTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2025) (SEQ. ID NO:2036)
          5'-CGTCBTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2026) (SEQ. ID NO:2037)
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5'-GTCBTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2027) (SEQ. ID NO:2038)
      5'-TCBTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2028) (SEQ. ID NO:2039)
      5'-CBTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2029) (SEQ. ID NO:2040)
      5'-BTGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2030) (SEQ. ID NO:2041)
       5'-TGGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2031) (SEQ. ID NO:2042)
       5'-GGCGGCGTCGGGCCGGGC-3' (FRAG. NO:2032) (SEQ. ID NO:2043)
       5'-GCGGCGTCGGGCCGGGC-3' (FRAG. NO:2033) (SEQ. ID NO:2044)
       5'-CGGCGTCGGGCCGGGC-3' (FRAG. NO:2034) (SEQ. ID NO:2045)
       5'-GGCGTCGGGCCGGGC-3' (FRAG. NO:2035) (SEQ. ID NO:2046)
       5'-GCGTCGGGCCGGGC-3' (FRAG. NO:2036) (SEQ. 1D NO:2047)
10
       5'-CGTCGGGCCGGGC-3' (FRAG. NO:2037) (SEQ. ID NO:2048)
       5'-GTCGGGCCGGGC-3' (FRAG. NO:2038) (SEQ. ID NO:2049)
       5'-TCGGGCCGGGC-3' (FRAG. NO:2039) (SEQ. ID NO:2050)
       5'-CGGGCCGGGC-3' (FRAG. NO:2040) (SEQ. ID NO:2051)
       5'-CCGCBGGCCBGGCGCCGCCGGCCGGCCGGCCG-3' (FRAG. NO:2041) (SEQ. ID NO:2052)
15
       5'-CCGCBGGCCBGGCCGCCGCCGGCCGGCC-3' (FRAG. NO:2042) (SEQ. ID NO:2053)
       5'-CCGCBGGCCBGGCGCCGCCGGCCGGCC3' (FRAG. NO:2043) (SEQ. ID NO:2054)
       5'-CCGCBGGCCBGGCCGCCGCCGGCCGGC-3' (FRAG. NO:2044) (SEQ. ID NO:2055)
       5'-CCGCBGGCCBGGCGCCGCCGCCGGCCGG-3' (FRAG. NO:2045) (SEQ. ID NO:2056)
       5'-CCGCBGGCCBGGCGCCGCCGCCGGCCG-3' (FRAG. NO:2046) (SEQ. ID NO:2057)
20
        5'-CCGCBGGCCBGGCGCCGCCGCCGCC-3' (FRAG. NO:2047) (SEQ. ID NO:2058)
        5'-CCGCBGGCCBGGCGCCGCCGCC3' (FRAG. NO:2048) (SEQ. 1D NO:2059)
        5'-CCGCBGGCCBGGCGCCGCCGC-3' (FRAG. NO:2049) (SEQ. ID NO:2060)
        5'-CCGCBGGCCBGGCGCCGCCG-3' (FRAG. NO:2050) (SEQ. ID NO:2061)
        5'-CCGCBGGCCBGGCGCCGCC-3' (FRAG. NO:2051) (SEQ. ID NO:2062)
 25
        5'-CCGCBGGCCBGGCGCGCCGC-3' (FRAG. NO:2052) (SEQ. ID NO:2063)
        5'-CCGCBGGCCBGGCGCGCCG-3' (FRAG. NO:2053) (SEQ. ID NO:2064)
        5'-CCGCBGGCCBGGCGCCC-3' (FRAG. NO:2054) (SEQ. ID NO:2065)
        5'-CCGCBGGCCBGGCGCGC-3' (FRAG. NO:2055) (SEQ. ID NO:2066)
        5'-CCGCBGGCCBGGCGCG-3' (FRAG. NO:2056) (SEQ. ID NO:2067)
 30
        5'-CCGCBGGCCBGGCGC-3' (FRAG. NO:2057) (SEQ. ID NO:2068)
        5'-CCGCBGGCCBGGCG-3' (FRAG. NO:2058) (SEQ. ID NO:2069)
        5'-CCGCBGGCCBGGGC-3' (FRAG. NO:2059) (SEQ. ID NO:2070)
         5'-CCGCBGGCCBGGG-3' (FRAG. NO:2060) (SEQ. ID NO:2071)
         5'-CCGCBGGCCBGG-3' (FRAG. NO:2061) (SEQ. ID NO:2072)
 35
         5'-CCGCBGGCCBG-3' (FRAG. NO:2062) (SEQ. ID NO:2073)
         5'-CCGCBGGCCB-3' (FRAG. NO:2063) (SEQ. ID NO:2074)
         5'-CCGCBGGCC-3' (FRAG. NO:2064) (SEQ. ID NO:2075)
         5'-CGCBGGCCBGGCGCCGCCGGCCGGCCGGCCGGCCG-3' (FRAG. NO:2065) (SEQ. ID NO:2076)
         5'-GCBGGCCBGGCGCCGCCGGCCGGCCGGCCG-3' (FRAG. NO:2066) (SEQ. ID NO:2077)
 40
         5'-CBGGCCBGGCGCGCCGCCGGCCGGCCG-3' (FRAG. NO:2067) (SEQ. ID NO:2078)
         5'-BGGCCBGGGCGCCGCCGGCCGGCCGGCCG-3' (FRAG. NO:2068) (SEQ. ID NO:2079)
         5'-GGCCBGGGCGCCGCCGGCCGGCCG-3' (FRAG. NO:2069) (SEQ. 1D NO:2080)
         5'-GCCBGGGCGCCGCCGGCCGGCCGGCCG-3' (FRAG. NO:2070) (SEQ. ID NO:2081)
         5'-CCBGGGCGCGCCGGCCGGCCGGCCG-3' (FRAG. NO:2071) (SEQ. ID NO:2082)
  45
         5'-CBGGGCGCGCCGGCCGGCCGGCCG-3' (FRAG. NO:2072) (SEQ. ID NO:2083)
          5'-BGGGCGCGCCGCCGGCCGGCCG-3' (FRAG. NO:2073) (SEQ. ID NO:2084)
         5'-GGGCGCCGCCGGCCGGCCG-3' (FRAG. NO:2074) (SEQ. ID NO:2085)
          5'-GGCGCGCCGCCGGCCGGCCG-3' (FRAG. NO:2075) (SEQ. ID NO:2086)
          5'-GCGCGCCGCCGGCCGGCCG-3' (FRAG. NO:2076) (SEQ. ID NO:2087)
  50
          5'-CGCGCCGCCGGCCGGGCCG-3' (FRAG. NO:2077) (SEQ. ID NO:2088)
          5'-GCGCCGCCGGCCGGCCG-3' (FRAG. NO:2078) (SEQ. ID NO:2089)
          5'-CGCCGCCGGCCGGCCG-3' (FRAG. NO:2079) (SEQ. ID NO:2090)
          5'-GCCGCCGGCCGGCCG-3' (FRAG. NO:2080) (SEQ. ID NO:2091)
          5'-CCGCCGGCCGGCCG-3' (FRAG. NO:2081) (SEQ. ID NO:2092)
   55
          5'-CGCCGGCCGGCCG-3' (FRAG. NO:2082) (SEQ. ID NO:2093)
          5'-GCCGGCCGGCCG-3' (FRAG. NO:2083) (SEQ. ID NO:2094)
          5'-CCGGCCGGGCCG-3' (FRAG. NO:2084) (SEQ. ID NO:2095)
          5'-CGGCCGGGCCG-3' (FRAG. NO:2085) (SEQ. ID NO:2096)
           5'-GGCCGGGCCG-3' (FRAG. NO:2086) (SEQ. ID NO:2097)
   60
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5'-GGGCGCBGGCTCCGCB-3' (FRAG. NO:2087) (SEQ. ID NO:2098)
      5'-GGGCCCCTGGCTCGGCCCGGGCCCGGCCCGGCCC-3' (FRAG. NO:2090) (SEQ. ID NO:2101)
      5'-GGGCCCCTGGCTCGGCCCGGGCCCGGCTTGCCCGCCCGGCC-3' (FRAG. NO:2091) (SEQ. ID NO:2102)
      5'-GGGCCCCTGGCTCGGCCCGCGGCCCGGCTTGCCCGGCC-3' (FRAG. NO:2092) (SEQ. ID NO:2103)
      5'-GGGCCCCTGGCTCGGCCCGGGCCCGGCTTGCCCGCCCGG-3' (FRAG. NO:2093) (SEQ. ID NO:2104)
      5'-GGGCCCCTGGCTCGGCCCGGGCCCGGCTTGCCCGCCC-3' (FRAG. NO:2095) (SEQ. ID NO:2106)
      5'-GGGCCCCTGGCTCGGCCCGCGGCCCGGCTTGCCCGCC-3' (FRAG. NO:2096) (SEQ. ID NO:2107)
10
      5'-GGGCCCCTGGCTCGGCCCGCGGCCCGGCTTGCCCGC-3' (FRAG. NO:2097) (SEQ. ID NO:2108)
      5'-GGGCCCCTGGCTCGGCCCCGCGGCCCGGCTTGCCCG-3' (FRAG. NO:2098) (SEQ. ID NO:2109)
      5'-GGGCCCCTGGCTCGGCCCGGGCCCGGCTTGCCC-3' (FRAG. NO:2099) (SEQ. ID NO:2110)
      5'-GGGCCCCTGGCTCGGCCCCGCGGCCCGGCTTGCC-3' (FRAG. NO:2100) (SEQ. ID NO:2111)
      5'-GGGCCCCTGGCTCGGCCCCGCGGCCCGGCTTGC-3' (FRAG. NO:2101) (SEQ. ID NO:2112)
15
      5'-GGGCCCCTGGCTCGGCCCCGCGGCCCGGCTTG-3' (FRAG. NO:2102) (SEQ. ID NO:2113)
      5'-GGGCCCCTGGCTCGGCCCGCGGCCCGGCTT-3' (FRAG. NO:2103) (SEQ. ID NO:2114)
      5'-GGGCCCCTGGCTCGGCCCCGCGGCCCGGCT-3' (FRAG. NO:2104) (SEQ. ID NO:2115)
      5'-GGGCCCCTGGCTCGGCCCGCGGCCCGGC-3' (FRAG. NO:2105) (SEQ. ID NO:2116)
      5'-GGGCCCCTGGCTCGGCCCGGGCCCGG-3' (FRAG. NO:2106) (SEQ. ID NO:2117)
20
      5'-GGGCCCCTGGCTCGGCCCGCGGCCCG-3' (FRAG. NO:2107) (SEQ. ID NO:2118)
      5'-GGGCCCCTGGCTCGGCCCCGCGGCCC-3' (FRAG. NO:2108) (SEQ. ID NO:2119)
      5'-GGGCCCCTGGCTCGGCCCCGCGGCC-3' (FRAG. NO:2109) (SEQ. ID NO:2120)
       5'-GGGCCCCTGGCTCGGCCCCGCGGC-3' (FRAG. NO:2110) (SEQ. ID NO:2121)
       5'-GGGCCCCTGGCTCGGCCCCGCGG-3' (FRAG. NO:2111) (SEQ. ID NO:2122)
25
       5'-GGGCCCCTGGCTCGGCCCCGCG-3' (FRAG. NO:2112) (SEQ. ID NO:2123)
       5'-GGGCCCCTGGCTCGGCCCCGC-3' (FRAG. NO:2113) (SEQ. ID NO:2124)
       5'-GGGCCCCTGGCTCGGCCCCG-3' (FRAG. NO:2114) (SEQ. ID NO:2125)
       5'-GGGCCCCTGGCTCGGCCCC-3' (FRAG. NO:2115) (SEQ. ID NO:2126)
       5'-GGGCCCCTGGCTCGGCCC-3' (FRAG. NO:2116) (SEQ. ID NO:2127)
30
       5'-GGGCCCCTGGCTCGGCC-3' (FRAG. NO:2117) (SEQ. ID NO:2128)
       5'-GGGCCCCTGGCTCGGC-3' (FRAG. NO:2118) (SEQ. ID NO:2129)
       5'-GGGCCCCTGGCTCGG-3' (FRAG. NO:2119) (SEQ. ID NO:2130)
       5'-GGGCCCCTGGCTCG-3' (FRAG. NO:2120) (SEQ. ID NO:2131)
       5'-GGGCCCCTGGCTC-3' (FRAG. NO:2121) (SEQ. ID NO:2132)
35
       5'-GGGCCCCTGGCT-3' (FRAG. NO:2122) (SEQ. ID NO:2133)
       5'-GCCCCTGGCTCGGCCCGGGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2124) (SEQ. ID NO:2135)
       5'-CCCTGGCTCGGCCCGGGCCCGGCCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2126) (SEQ. ID NO:2137)
 40
       5'-CCTGGCTCGGCCCGGGCCCGGCCTTGCCCGCCCGGCCCGG-3' (FRAG, NO:2127) (SEQ. ID NO:2138)
       5'-TGGCTCGGCCCGGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2129) (SEQ. ID NO:2140)
       5'-GGCTCGGCCCCGCGGCCCGGCCCGGCCCGGCCCGG-3' (FRAG. NO:2130) (SEQ. ID NO:2141)
       5'-GCTCGGCCCCGCGGCCCGGCCCGGCCCGGCCCGG-3' (FRAG. NO:2131) (SEQ. ID NO:2142)
 45
       5'-CTCGGCCCCGCGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2132) (SEQ. ID NO:2143)
       5'-TCGGCCCCGCGGCCCGGCCCGGCCCGG-3' (FRAG. NO:2133) (SEQ. ID NO:2144)
        5'-CGGCCCCGCGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2134) (SEQ. ID NO:2145)
       5'-GGCCCCGCGGCCCGGCCCGGCCCGGCCCGG-3' (FRAG. NO:2135) (SEQ. ID NO:2146)
        5'-GCCCCGCGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2136) (SEQ. ID NO:2147)
 50
        5'-CCCCGCGGCCCGGCCCGGCCCGG-3' (FRAG, NO:2137) (SEQ. ID NO:2148)
        5'-CCCGCGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2138) (SEQ. ID NO:2149)
        5'-CCGCGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2139) (SEQ. ID NO:2150)
        5'-CGCGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2140) (SEQ. ID NO:2151)
        5'-GCGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2141) (SEQ. ID NO:2152)
 55
        5'-CGGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2142) (SEQ. ID NO:2153)
        5'-GGCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2143) (SEQ. ID NO:2154)
        5'-GCCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2144) (SEQ. ID NO:2155)
        5'-CCCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2145) (SEQ. ID NO:2156)
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5'-CCGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2146) (SEQ. 1D NO:2157)
       5'-CGGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2147) (SEQ. ID NO:2158)
       5'-GGCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2148) (SEQ. ID NO:2159)
       5'-GCTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2149) (SEQ. ID NO:2160)
       5'-CTTGCCCGCCCGGCCCGG-3' (FRAG. NO:2150) (SEQ. ID NO:2161)
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       5'-TTGCCCGCCCGGCCCGG-3' (FRAG. NO:2151) (SEQ. ID NO:2162)
       5'-TGCCCGCCCGGCCCGG-3' (FRAG. NO:2152) (SEQ. ID NO:2163)
       5'-GCCCGCCCGGCCCGG-3' (FRAG. NO:2153) (SEQ. ID NO:2164)
       5'-CCCGCCCGGCCCGG-3' (FRAG. NO:2154) (SEQ. ID NO:2165)
       5'-CCGCCCGGCCCGG-3' (FRAG. NO:2155) (SEQ. ID NO:2166)
10
       5'-CGCCCGGCCCGG-3' (FRAG. NO:2156) (SEQ. ID NO:2167)
       5'-GCCCGGCCCGG-3' (FRAG. NO:2157) (SEQ. ID NO:2168)
       5'-GGCGGGGGGGGGGGCGCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2158) (SEQ. ID NO:2169)
       5'-GGCGGGGGCGCCCCTGGCTCGCCTBGGGCCC-3' (FRAG. NO:2159) (SEQ. ID NO:2170)
       5'-GGCGGGGGGGGGCGCCTGGCCTBGGGCC-3' (FRAG. NO:2160) (SEQ. ID NO:2171)
15
       5'-GGCGGGGGCGCCCCGCCTBGGGC-3' (FRAG. NO:2161) (SEQ. ID NO:2172)
       5'-GGCGGGGGGGGGCGCCTGGCTCGCCTBGGG-3' (FRAG. NO:2162) (SEQ. ID NO:2173)
        5'-GGCGGGGGCGGCGCCTGGCTCGCCTBGG-3' (FRAG. NO:2163) (SEQ. ID NO:2174)
        5'-GGCGGGGGGGGGGCGCCTGGCTCGCCTBG-3' (FRAG. NO:2164) (SEQ. ID NO:2175)
        5'-GGCGGGGGGGGGGGCGCCTGGCTCGCCTB-3' (FRAG. NO:2165) (SEQ. ID NO:2176)
20
        5'-GGCGGGGGCGCCGCCTGGCTCGCCT-3' (FRAG. NO:2166) (SEQ. ID NO:2177)
        5'-GGCGGGGGGGGGGCGCCTGGCTCGCC-3' (FRAG. NO:2167) (SEQ. ID NO:2178)
        5'-GGCGGGGGGGGGCGCCTGGCTCGC-3' (FRAG. NO:2168) (SEQ. ID NO:2179)
        5'-GGCGGGGGCGCGCCCTGGCTCG-3' (FRAG. NO:2169) (SEQ. ID NO:2180)
        5'-GGCGGGGGGGGGCGCCTGGCTC-3' (FRAG. NO:2170) (SEQ. ID NO:2181)
25
        5'-GGCGGGGGCGCCCTGGCT-3' (FRAG. NO:2171) (SEQ. ID NO:2082)
        5'-GGCGGGGGGGGGCGCCTGGC-3' (FRAG. NO:2172) (SEQ. ID NO:2183)
        5'-GGCGGGGGCGCCGCCTGG-3' (FRAG. NO:2173) (SEQ. ID NO:2184)
        5'-GGCGGGGGGGGGGCGCCTG-3' (FRAG. NO:2174) (SEQ. ID NO:2185)
        5'-GGCGGGGGGGGGGCGCCT-3' (FRAG. NO:2175) (SEQ. ID NO:2186)
30
        5'-GGCGGGGGCGCCGCC-3' (FRAG. NO:2176) (SEQ. ID NO:2187)
        5'-GGCGGGGGCGCGCGCGC-3' (FRAG. NO:2177) (SEQ. ID NO:2188)
        5'-GGCGGGGGCGCGCG-3' (FRAG. NO:2178) (SEQ. ID NO:2189)
        5'-GGCGGGGGCGGCGC-3' (FRAG. NO:2179) (SEQ. ID NO:2190)
        5'-GGCGGGGGCGCGG-3' (FRAG. NO:2180) (SEQ. ID NO:2191)
 35
        5'-GGCGGGGGCGCG-3' (FRAG. NO:2181) (SEQ. ID NO:2192)
        5'-GGCGGGGGCGC-3' (FRAG. NO:2182) (SEQ. ID NO:2193)
        5'-GGCGGGGGCGG-3' (FRAG. NO:2183) (SEQ. ID NO:2194)
        5'-GCGGGGGCGCCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2184) (SEQ. ID NO:2195)
        5'-CGGGGGCGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2185) (SEQ. ID NO:2196)
 40
        5'-GGGGGCGGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2186) (SEQ. ID NO:2197)
        5'-GGGGCGGCGCCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2187) (SEQ. ID NO:2198)
         5'-GGGCGGCGGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG, NO:2188) (SEQ. 1D NO:2199)
         5'-GGCGGCGCCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2189) (SEQ. 1D NO:2200)
         5'-GCGGCGGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2190) (SEQ. ID NO:2201)
 45
         5'-CGGCGGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2191) (SEQ. ID NO:2202)
         5'-GGCGGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2192) (SEQ. ID NO:2203)
         5'-GCGGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2193) (SEQ. ID NO:2204)
         5'-CGGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2194) (SEQ. ID NO:2205)
         5'-GGCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2195) (SEQ. ID NO:2206)
 50
         5'-GCGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2196) (SEQ. ID NO:2207)
         5'-CGCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2197) (SEQ. ID NO:2208)
         5'-GCCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2198) (SEQ. ID NO:2209)
         5'-CCTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2199) (SEQ. ID NO:2210)
         5'-CTGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2200) (SEQ. ID NO:2211)
 55
         5'-TGGCTCGCCTBGGGCCCC-3' (FRAG. NO:2201) (SEQ. ID NO:2212)
         5'-GGCTCGCCTBGGGCCCC-3' (FRAG. NO:2202) (SEQ. ID NO:2213)
         5'-GCTCGCCTBGGGCCCC-3' (FRAG. NO:2203) (SEQ. ID NO:2214)
         5'-CTCGCCTBGGGCCCC-3' (FRAG. NO:2204) (SEQ. 1D NO:2215)
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5'-TCGCCTBGGGCCCC-3' (FRAG. NO:2205) (SEQ. ID NO:2216)
       5'-CGCCTBGGGCCCC-3' (FRAG. NO:2206) (SEQ. ID NO:2217)
       5'-GCCTBGGGCCCC-3' (FRAG. NO:2207) (SEQ. ID NO:2218)
       5'-CCTBGGGCCCC-3' (FRAG. NO:2208) (SEQ. ID NO:2219)
       5'-CTBGGGCCCC-3' (FRAG. NO:2209) (SEQ. ID NO:2220)
       5'-GGGTGGGCBCGGCGGCC-3' (FRAG. NO:2210) (SEQ. ID NO:2221)
       5'-GGTCGGCGBBGBGCTCGTCGTGGC-3' (FRAG. NO:2211) (SEQ. ID NO:2222)
       5'-GGTCGGCGBBGBGCTCGTCGTGG-3' (FRAG. NO:2212) (SEQ. ID NO:2223)
       5'-GGTCGGCGBBGBGCTCGTCGTG-3' (FRAG. NO:2213) (SEQ. ID NO:2224)
       5'-GGTCGGCGBBGBGCTCGTCGT-3' (FRAG. NO:2214) (SEQ. ID NO:2225)
10
       5'-GGTCGGCGBBGBGCTCGTCG-3' (FRAG. NO:2215) (SEQ. ID NO:2226)
       5'-GGTCGGCGBBGBGCTCGTC-3' (FRAG. NO:2216) (SEQ. ID NO:2227)
       5'-GGTCGGCGBBGBGCTCGT-3' (FRAG. NO:2217) (SEQ. ID NO:2228)
       5'-GGTCGGCGBBGBGCTCG-3' (FRAG. NO:2218) (SEQ. ID NO:2229)
       5'-GGTCGGCGBBGBGCTC-3' (FRAG. NO:2219) (SEQ. ID NO:2230)
15
       5'-GGTCGGCGBBGBGCT-3' (FRAG. NO:2220) (SEQ. ID NO:2231)
       5'-GGTCGGCGBBGBGC-3' (FRAG. NO:2221) (SEQ. ID NO:2232)
        5'-GGTCGGCGBBGBG-3' (FRAG. NO:2222) (SEQ. ID NO:2233)
        5'-GGTCGGCGBBGB-3' (FRAG. NO:2223) (SEQ. ID NO:2234)
        5'-GGTCGGCGBBG-3' (FRAG. NO:2224) (SEQ. 1D NO:2235)
20
        5'-GTCGGCGBBGBGCTCGTCGTGGC-3' (FRAG. NO:2225) (SEQ. ID NO:2236)
        5'-TCGGCGBBGBGCTCGTCGTGGC-3' (FRAG. NO:2226) (SEQ. ID NO:2237)
        5'-CGGCGBBGBGCTCGTCGTGGC-3' (FRAG. NO:2227) (SEQ. ID NO:2238)
        5'-GGCGBBGBGCTCGTCGTGGC-3' (FRAG. NO:2228) (SEQ. ID NO:2239)
        5'-GCGBBGBGCTCGTCGTGGC-3' (FRAG. NO:2229) (SEQ. ID NO:2240)
25
        5'-CGBBGBGCTCGTCGTGGC-3' (FRAG. NO:2230) (SEQ. ID NO:2241)
        5'-GBBGBGCTCGTCGTGGC-3' (FRAG. NO:2231) (SEQ. ID NO:2242)
        5'-BBGBGCTCGTCGTGGC-3' (FRAG. NO:2232) (SEQ. ID NO:2243)
        5'-BGBGCTCGTCGTGGC-3' (FRAG. NO:2233) (SEQ. 1D NO:2244)
        5'-GBGCTCGTCGTCGC-3' (FRAG. NO:2234) (SEQ. ID NO:2245)
30
        5'-BGCTCGTCGTGGC-3' (FRAG. NO:2235) (SEQ. ID NO:2246)
        5'-GCTCGTCGTGGC-3' (FRAG. NO:2236) (SEQ. ID NO:2247)
        5'-CTCGTCGTGGC-3' (FRAG. NO:2237) (SEQ. ID NO:2248)
        5'-TCGTCGTGGC-3' (FRAG. NO:2238) (SEQ. ID NO:2249)
        5'-GGGGCCCCGCCGCCCGCC-3' (FRAG. NO:2239) (SEQ. ID NO:2250)
35
        5'-GGGGCCCCGCGCCCCCC-3' (FRAG. NO:2240) (SEQ. 1D NO:2251)
        5'-GGGGCCCGCGCCGCCG-3' (FRAG. NO:2241) (SEQ. ID NO:2252)
        5'-GGGGCCCCGCCCCC-3' (FRAG. NO:2242) (SEQ. ID NO:2253)
        5'-GGGGCCCCGCGCCCCC'3' (FRAG. NO:2243) (SEQ. ID NO:2254)
        5'-GGGGCCCCGCGCCGC-3' (FRAG. NO:2244) (SEQ. ID NO:2255)
 40
         5'-GGGGCCCCGCGCCG-3' (FRAG. NO:2245) (SEQ. ID NO:2256)
         5'-GGGGCCCCGCGCC-3' (FRAG. NO:2246) (SEQ. ID NO:2257)
         5'-GGGGCCCCGCGC-3' (FRAG. NO:2247) (SEQ. ID NO:2258)
         5'-GGGCCCGCGCCGCCCGCC-3' (FRAG. NO:2248) (SEQ. ID NO:2259)
         5'-GGCCCCGCCCCCCCCC3' (FRAG. NO:2249) (SEQ. ID NO:2260)
 45
         5'-GCCCGCGCCCCCCCC3' (FRAG. NO:2250) (SEQ. ID NO:2261)
         5'-CCCCGCGCCCCCCCC-3' (FRAG. NO:2251) (SEQ. ID NO:2262)
         5'-CCCGCGCCCCCCCC-3' (FRAG. NO:2252) (SEQ. ID NO:2263)
         5'-CCGCGCCCCCCC-3' (FRAG. NO:2253) (SEQ. ID NO:2264)
         5'-CGCGCCGCCCGCC-3' (FRAG. NO:2254) (SEQ. ID NO:2265)
 50
         5'-GCGCCGCCCGCC-3' (FRAG. NO:2255) (SEQ. ID NO:2266)
         5'-CGCCGCCCGCC-3' (FRAG. NO:2256) (SEQ. ID NO:2267)
         5'-GCCGCCCGCC-3' (FRAG. NO:2257) (SEQ. ID NO:2268)
         5'-GGGGCGCGCGGGGCCGCCGGG-3' (FRAG. NO:2258) (SEQ. 1D NO:2269)
         5'-GGCGGGGBGCGGCCCGGGCCC-3' (FRAG. NO:2259) (SEQ. ID NO:2270)
  55
         5'-GGCGCGTCGCCCTCGCCCCBGTCGGGCTCGCGC-3' (FRAG. NO:2260) (SEQ. ID NO:2271)
         5'-GCGCGGGCBBCBGCGBGCCGGGCGCG-3'.(FRAG. NO:2261) (SEQ. ID NO:2272)
         5'-GCGCBCGGGCCCBCCTGCGCGGGC-3' (FRAG. NO:2262) (SEQ. ID NO:2273)
         5'-GGGCGGGGTGGGCTGCCCTGCGGCCGCC-3' (FRAG, NO:2263) (SEQ. ID NO:2274)
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5'-GGGCTGCTGCGCGGCGGCTCCGGCGA-3' (FRAG. NO:2264) (SEQ. ID NO:2275)
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- 5'-CTCCCGGGCGGGGCCGGGGCGCGGGG-3' (FRAG, NO:2265) (SEQ. ID NO:2276)
- 5'-GGGCTGCCGCGGTCCGGGCCCCTCTTGCCGGCG-3' (FRAG. NO:2266) (SEQ. 1D NO:2277)
- 5'-GCGCTCGCCGCCGCTGCCGG-3' (FRAG. NO:2267) (SEQ. ID NO:2278)
- 5'-GCGCCGCTTGGCCTTGTCGCGGC-3' (FRAG. NO:2268) (SEQ. ID NO:2279)
 - 5'-GCTGCTCCBCGCGCTGG-3' (FRAG. NO:2269) (SEQ. ID NO:2280)
 - 5'-GCCGGBGGCCGGCCBGGTCCCGCG-3' (FRAG. NO:2270) (SEQ. ID NO:2281)
 - 5'-CCCGGCGGCCGGCBGGBBGGGCGGGCTGGGC-3' (FRAG. NO:2271) (SEQ. ID NO:2282)
 - 5'-GTCTCTCCCGCCCCGGCCGCGCGCG-3' (FRAG. NO:2272) (SEQ. ID NO:2283)
- 10 5'-GGGCGTCCGGTCCGGGCCGTCGGG-3' (FRAG. NO:2273) (SEQ. ID NO:2284)
 - 5'-GCGGGCACGCGGCGCTCTGGCGTCGGC-3' (FRAG. NO:2274) (SEQ. ID NO:2285)

Where B is adenosine, or, more preferbly, replaces adenosine and is a universal base, and adenosine A2a receptor agonist, an adenosine A2b receptor antagonist, an adenosine A3 receptor antagonist, or an adenosine A1 receptor antagonist

15 Bradykinin Receptor Anti-sense Oligonucleotide Fragments

- 5'-GGTGBCBTTG BGCBTGTCGG CGCGGTCCCG TTBBGBGTGG GCCCGCCAGC CCAGCCACTC CACTTGGGGG CGGGTGGCCA GCACGAACAG CACCCAGAGG AAGGGGGGCG GCCCAGAAGG GCAGCCCGCA GGCCAGGATC AGGTCTGCTG CGGCCGGAGA TAATGGCATT CACCACGCGG CGGCCCAGCG CACGCCGCG ATCCGGCCCG GGTTCTGACC TGCAGCCCCC GTCTCCTTGG CATTCCTGGG CCCCAGTCAC TCCTCTCCCT GCCCCCCTTG CTGGGGCAGG GACGGGGTG BCBTTGBGCB TGTCGGCGCG
- 20 GTCCCGTTBB GBGTGGCCC GCCAGCCCAG CCACTCCACT TGGGGGCGGG TGGCCAGCAC GAACAGCACC CAGAGGAAGG
 GGGGCGGCCC AGAAGGCCAG CCCGCAGGCC AGGATCAGGT CTGCTGCGGC CGGAGATAAT GGCATTCACC ACGCGGCGC
 CCAGCGCACG CCGCGCATCC GGCCCGGGTT CTGACCTGCA GCCCCCGTCT CCTTGGCATT CCTGGGCCCC AGTCACTCCT
 CTCCCTGCCC CCCTTGCTGG GGCAGGGACG GCCGTGTTGT CBGTGGTGCT GCCCGTTTGB GGTBTGGCGC TCCBCCBBTT
 CCCTTTTCTC CTTGTTTTCC GTTTCTCTTG CCGTCTGTGG TT-3' (FRAG. NO:2275) (SEQ. ID NO:2286)
- 25 5'-GGTGBCBTTGBGCBTGTCGGCGC-3' (FRAG. NO:2276) (SEQ. ID NO:2287)
 - 5'-GGTCCCGTTBBGBGTGGGCCC-3' (FRAG. NO:2277) (SEQ. ID NO:2288)
 - 5'-GCCAGCCCAGCCACTCCACTTGGGGGC-3' (FRAG. NO:2278) (SEQ. ID NO:2289)
 - 5'-GGGTGGCCAGCACGAACAGCACCCAGAGGAAGGGGGGC-3' (FRAG. NO:2279) (SEQ. ID NO:2290)
 - 5'-GGCCCAGAAGGGCAGCCCGCAGGCCAGGATCAGGTCTGCTGCGGCC-3' (FRAG. NO:2280) (SEQ. ID NO:2291)
- 30 5'-GGAGATAATGGCATTCACCACGCGGC-3' (FRAG. NO:2281) (SEQ. ID NO:2292)
 - 5'-GGCCCAGCGCACGCCGCGCATCCGGCCC-3' (FRAG. NO:2282) (SEQ. ID NO:2293)
 - 5'-GGGTTCTGACCTGCAGCCCCC-3' (FRAG. NO:2283) (SEQ. ID NO:2294)
 - 5'-GTCTCCTTGGCATTCCTGGGCCC-3' (FRAG. NO:2284) (SEQ. ID NO:2295)
 - 5'-CAGTCACTCCCTGCCCCC-3' (FRAG. NO:2285) (SEQ. ID NO:2296)
- 35 5'-CTTGCTGGGGCAGGGACGG-3' (FRAG. NO:2286) (SEQ. ID NO:2297)
 - 5'-GGTGBCBTTGBGCBTGTCGGCGC-3' (FRAG. NO:2287) (SEQ. ID NO:2298)
 - 5'-GGTCCCGTTBBGBGTGGGCCC-3' (FRAG. NO:2288) (SEQ. ID NO:2299)
 - 5'-GCCAGCCCAGCCACTCCACTTGGGGGC-3' (FRAG. NO:2289) (SEQ. ID NO:2300) 5'-GGGTGGCCAGCACGAACAGCACCCAGAGGAAGGGGGGC-3' (FRAG. NO:2290) (SEQ. ID NO:2301)
- 40 5'-GGCCCAGAAGGGCAGCCCGCAGGCCAGGATCAGGTCTGCTGCGGCC-3' (FRAG. NO:2291) (SEQ. 1D NO:2302)
 - 5'-GGAGATAATGGCATTCACCACGCGGC-3' (FRAG. NO:2292) (SEQ. ID NO:2303)
 - 5'-GGCCCAGCGCACGCCGCGCATCCGGCCC-3' (FRAG. NO:2293) (SEQ. ID NO:2304)
 - 5'-GGGTTCTGACCTGCAGCCCCC-3' (FRAG. NO:2294) (SEQ. ID NO:2305)
 - 5'-GTCTCCTTGGCATTCCTGGGCCC-3' (FRAG, NO:2295) (SEQ. ID NO:2306)
- 45 5'-CAGTCACTCCTCTCCCTGCCCCC-3' (FRAG. NO:2296) (SEQ. ID NO:2307)
 - 5'-CTTGCTGGGGCAGGGACGG-3' (FRAG. NO:2297) (SEQ. ID NO:2308)
 - 5'-CCGTGTTGTCBGTGGTGCTG-3' (FRAG. NO:2298) (SEQ. ID NO:2309)
 - 5'-CCCGTTTGBGGTBTGGC-3' (FRAG. NO:2299) (SEQ. ID NO:2310)
 - 5'-GCTCCBCCBBTTCCCTTTTCTCC-3' (FRAG. NO:2300) (SEQ. ID NO:2311)
- 50 5'-TTGTTTTCCGTTTCTCTTG-3' (FRAG. NO:2301) (SEQ. ID NO:2312)
 - 5'-CCGTCTGTGGTT-3' (FRAG. NO:2302) (SEQ. ID NO:2313)

In one preferred embodiment, the links between neighboring mononucleotides are

phosphodiester links. In another preferred, at least one mononucleotide phosphodiester residue of the anti-sense oligonucleotide(s) is substituted by a methylphosphonate, phosphotriester, phosphorothioate, phosphorodithioate, boranophosphate, formacetal, thioformacetal, thioether, carbonate, carbamate, sulfate, sulfonate, sulfamate, sulfonamide, sulfone, sulfite, sulfoxide, sulfide, hydroxylamine, 2'-O-methyl, methylene(methyimino), methyleneoxy (methylimino), phosphoramidate residues, and combinations thereof. The MTA oligos having one or more phosphodiester residues substituted by one or more of the other residues are generally longer lasting, given that these residues are more resistant to hydrolysis than the phosphodiester residue. In some cases up to about 10%, about 30%, about 50%, about 75%, and even all phosphodiester residues may be substituted (100%).

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Typically, the multiple target anti-sense oligonucleotide (MTA oligo) of the invention comprises at least about 7 mononucleotides, in some instances up to 60 and more mononucleotides, preferably about 10 to about 36, and more preferably about 12 to about 21 mononucleotides. However, other lengths are also suitable depending on the length of the target macromolecule. Examples of the MTA oligos of the invention are provided in Table 3 below, which includes ninety-four sequences (SEQ ID NOS.: 2317 through 2411).

Table 3: MTA Oligos, Location Targeted & Target

ITA Oligo	No.	SEQ. ID Location Compound Targe Targeted	t -
UMNFKBP65A AS			
CC GGC CCC GCC TCG TGC C GT CCB TGC CGC GGG CCC CC CCG CTG CTT GGG CTG CTC	2317 2318 2319		
CT GTG CTC CTC TCG CCT GGG GG TGG GGT TCT TGG TGG TC TCC CTG GTC CTG TG	20123 22223 223324 223324	5' = 137 EPI 2195 5'=159 EPI 2196 5'=166 EPI 2197 5'=362 EPI 2198 5'=401 EPI 2198	
ביים ביים ביים	UMNFKBP65A AS CC GGC CCC GCC TCG TGC C ET CCB TGC CGC GGG CCC CCC CTG CTT GGG CTG CTC CGC CGG G ET GTG CTC CTC TCG CCT GGG ET GTG CTC CTC TCG TGG ET GTG CTC CTC TCG TGG ET GTG CTC CTC TGG ET GTG CTC CTC TGG ET GTC CTG GTC CTG TG	UMNFKBP65A AS CC GCC CCC GCC GGG CCC CTC CCC CCC CTC CTC CTC CCC CTC CTC	No. Targeted UMNFKBP65A AS CC GGC CCC GCC TCG TGC C 2317 5, = 1 EPI 2192 ET CCB TGC CGC GGG CCC 2318 5, = 28 (AUG) EPI 2193 CC CCG CTG CTT GGG CTC 2319 5, = 65 EPI 2194 ET GTG CTC CTC TCG CCT GGG 2320 5, = 137 EPI 2196 ET GTG CTC CTC TGG TGG 2322 5, = 136 EPI 2197 ET GTG CTG GTC CTG TGG 2322 5, = 136 EPI 2197

Table 3: MTA Oligos, Location Targeted & Target (Cont'd)

MTA Oligo		SEQ. ID Location Compound Target
	No.	Targeted
GCC TCG GGC CTC CC GGC TGG GGT CTG CGT GGC CGG GGG TCG GTG GGT CCG CTG	2326 2327	5'=697 EPI 2201 5'=769 EPI 2202
CCG CTG GGG CTT GGG CTT GGG C GGG CTT GGG GCC TGG GCC GCC TGG GTG GGC TTG GGG GCT GGG TCT GTG CTG TTG CC GCT GGG TCT GTG CTG TTG CC GTT GTG TGG GGG GCC GTT GTG TGG	2331 2332 2333 2333	5'=953 EPI 2203 5'=1022 EPI 2204 5'=1208 EPI 2205 5'=1272 EPI 2206 5'=1362 EPI 2207 5'= 1451 EPI 2208
GCT GGG TTG GGG GGC CTC TGG GCT GTC GCC CCG GGG CCC CC TGG CTC CCC CCT CC GCT CCC CCC TTT CC GCT CCC CCC TTT CC CGG ACG AAG AGG A GGC TTT GTG GGC TC GCC TGC TCT CCC CC	23333333333333333333333333333333333333	5'=1511 EPI 2209 5'=1550 EPI 2210 5'=1762 EPI 2211 5'=1863 EPI 2212 5'=1979 EPI 2213 5'=2011 EPI 2215
CCC GGC CCC GCC BCG BBC C	2341	intron EPI 2192-01A HSU50136 C4 Syntha:
CCC GGC CCC GCC BCG CCC GGC CCC GCC BCG BBC C CCC GGC CCC GCC BCG CCC GBC CCC GCC TCB BG	2342 2343 2344 2345	intron EPI 2192-01B 5'untr EPI 2192-02A HUMLIPOX 5 5'untr EPI 2192-02B trans EPI 2192-03A HSNFKBS
SUDUNIT CCC GBC CCC GCC TC CCG GCC CCG CCT C CCC GBB CCC GCB TBG TGC C	2346 2347 2348	trans EPI 2192-03B 5'untr EPI 2192-04 TGFBR1 5'trans EPI 2192-05A HSU58198 : enhan
CCC GCB TBG TGC C CCC GGB CCC BCC BBG TGC C CBG BBC CCG CCT CGT GCC C CCG CCT CGT GCC CCG GCB CCG CCT CBT GCC CCG GCC CCG CCB CBT GCC	2349 23551 23552 23553 2354	5'untr EPI 2192-05B 3'trans EPI 2192-06 HSVECAD intron EPI 2192-07A NFKB2 intron EPI 2192-07B NFKB2 5'trans EPI 2192-08 Carboxype 3'trans EPI 2192-09 HUMADRA2C 3'trans EPI 2192-09 G2adr kid
CCC GBC CCC GBC TCG CCC GGC CBC GBC TCG	2355 2356	5'untrs EPI 2192-10 HUMFK506B 5'untrs EPI 2192-11 HSNBARKS1 Badr kina 5'UTR EPI 2192-12 HSNFXN1
CCC GGC CCB GCC TBG (NFKB1) CCC GGC BCB GBC TCG TBC C	2357	
ccc GGC BCB GBC TCG TBC C (transcrp.	2358	3'UTR EPI 2192-13 HSILF
CCC GGC CCC GCC BCG Syn/5-LO MTA	2359	Factor IL EPI-2192-14 NFKB/C4 /TGFBrec1
CCC GGC CCC GCC BCG	2360	/5-1.0 MTA
TCC BTG CCG CGG GC TCC BTG CCB CGG GCC TCC BTG CCB CGG GCC TCC BTG CCB CBG GCC GTC CBT GBC GCG	2361 2362 2363 2364 2365 2366	3' trans EPI-2193-01 MET Oncog 3' trans EPI-2193-02 HSFGR2(IG mid cod EPI-2193-03 5-LO mid cod EPI-2193-04 HUMTK14 3'trans EPI-2193-05 HUMTNFR AUG
TC CBT GBC GCG GG	2300	Probl.HUMPTCH cardiac
K+channel TCT GBG CTC CTC TBB CCT GGG cytotox.	2367	
CTG TGC BCC TBB CBC CTG GG	2368	ser. prot intr EPI-2195-02 HSINOSX08 inducible
TGT GBT CCB CTB GBC TGG G	2369	EPI-2195-03 HUMACHRM2 musc. m2 acetyl
TCT GTB CTC BBC TCB CCT G	2370	EPI-2195-04 s86371s1 Neurokini rec
TGC TCC TCB CBB CTG GG	2371	

<u>Table 3</u>: MTA Oligos, Location Targeted & Target (Cont'd)

MTA	4 (Oli	go				No.	SEQ. ID Lo	ocation	Compound	Target
											inflam. factor
CTC	CI	C	TBG	CCT	GG		2372		EPI-21	.95-06	HSNBARKS4 β-adr rec kinase
GTG	CI	C	CBB	TCB	BCT	GGG	2373		EPI-21	.95-07	HSTNFR2SO TNF R2
GTG	CE	3C	CBB	TCB	CCT	GGG	2374		EPI-21	.95-08	humfkbp fk506 binding protein
TCT	G1	ľG	CBC	CTC	TBG	BCT	2375	exon	EPI-21	95-09	HSNBARKS1 adr. receptor
CTG TGT	TI	BB CT	TCC BBT	TBB CBC	CBC BCB	CTG G TGG G	2376 2377	intron	EPI-21		kinase HUMIL8 HSU50157 PDE4
GTG CTG	TO	ЗĊ	CBC BCC	TCT	CCT		2378 2379	3'UTR	EPI-22		IL-2 R IL-6 R HSIL6R
G TO CBC C C	GC C TC	BC FC TC	C BC TCB B CC	BCT CCT CCT GCC	SC CI GGG SG	CTG IG	2380 2381 2382 2383 2384	intr/ex intr/ex coding coding coding	EPI-22 EPI-22	203-06A 203-06B 203-07A 03-07B 203-08	HSIL2rG6 HSIL2rG6 HUMIL71 IL-7 HUMII IL-6 R HSI6REC
TGC	T	CC	TCB	CGC	C		2385	intron P	DGF A I	EPI-2303-09	chain
			GBT	CTG	G		2386	3'utr	EPI-2	199-01	HUMPDGFAE GATA-4 Factor fo IL-5
GGT GGT			BBT TGB	TGG TGB	TCT TCT	TGG G	2387 2388	Coding Far 5'UT		199-02 2199-03	TNFG HUMTH HSSUBPIG (Sub P r)
GGG	T	ТВ	BBG	TTG	BTC	TGG	2389	Coding		199-04	Neutrophi adh. R_HUMNARI
				TTG		TGG	2390	HSHM2		199-05	m2 muscarini R L1 leuk.
TTG	T	TG	TBG	BTC	TGG		2391	HUML1CAM	r FFT-2	199-00	adh.
GGG	T	CB BG CG	GBG GTG GBG		CBG C CBG	CTG	2392 2393 2394 2395	coding S71424S2 coding HUMITGF HUMNK65E	EPI-2 EPI-2 EPI-2	203-01 203-02 203-03 203-04 -2206-01	prot. HumgaTa21 IGE eps HSGCSFR2 TGFβ3 NFKB/NK Cell Activati
GGG	Т	GG	GCT	TGG	G		2396	HUMPEREE	B EPI	2206-02	Protein NFKB/ Prostagl EP3 Recep

Table 3: MTA Oligos, Location Targeted & Target (Cont'd)

MTA Oligo	No.	EQ. ID Location Compound Target Targeted
CCTGGGTGGGBBTGGG	2397	EPI 2206-03 HSNF2B/GCSF NFKB/ Granulocyte CSF/Transcr
CCTGGBTGGGCBTGGG	2398	Factor NF2E EPI-2206-04 HUMLAP/NFKE Leuk Prot.
GCCTGBGTGBBCTTGGG	2399	EPI 2206-05 NFKB/ Endotheline N2 S63833
CCCAVGVCCVCCCAGGC	2400	EPI 2206-06 NFKBAS13/B Lymph. Ser Prot. Kinas
AGCCCACCCAGGC	2401	EPI 2206-07 NFKBAS13/G HSGCSFR1
BCCTGGGTGGGCTB	2402	EPI 2206-08 NFKBAS13/G /NK7TCELLAG Prot.
GGTGGGCTTGGG	2403	EPI 2206-09 NFKBAS13 /HSTGFB1 TO
CCBBGGTGGGCTTGGG	2404	EPI 2206-10 NFKBAS13 /HSTGFB1 T
CTGGGTGGGBBTGGG	2405	EPI 2206-11 NFKBAS13/ HSGCSFR1 GCSFR1
CCBGGGTGGGCTTGG	2406	EPI 2206-12 NFKBAS13/ HUMCD30A Lymph.Act. Antig.(Cod
GGGTGGGCTTGG	2407	EPI-2206-12B NFKBAS13/
HUMCD30A CCTGBGTGBGCBTGGG	2408	EPI 2206-13 NFKBAS13/ HUMCAM1V Vasc. Endo Cell Adh. Molec
B: Universal Base	· · · · · · · · · · · · · · · · · · ·	

The MTA oligos of Table 3 are suitable for use with two or more of the targets listed in Table 4 below.

PCT/US98/19419 WO 99/13886

Targets for the MTA Oligos of Table 3 Table 4:

	Compound	Target	
	EPI 2010	Adenosine A1 receptor	
	EPI 2045	Adenosine A3 receptor	
5	EPI 2873, EPI 2193	NFxB	
•	EPI 1873	Interleukin-l	
	EPI 1857	Interleukin -5	
	EPI 2945	Interleukin -4	
	EPI 2977	Interleukin -8	
10	EPI 2031	5-Lipoxygenase	
••	EPI 1898	Leukotriene C-4 Synthase	
	EPI 1856	Eotaxin	
	EPI 1131	ICAM	
	EPI 1085	VCAM	
15	EPI 2085	TNFa	
	EPI 1908	PAF	
	EPI 1925	IL-4 receptor	
	EPI 2643	β2 aderenergic receptor kinase	
	EPI 2934	Tryptase	
20	EPI 2033	Major Basic Protein	
	EPI 2795	Eosinophil Peroxidase	

NfkB: nuclear factor kB

NIKE: nuclear factor KB
ICAM: intracellular adhesion molecule
VCAM: vascular cell adhesion molecule
TNF: tumor necrosis factor
PAF: platelet activating factor

In a most preferred embodiment for use in the lung, the MTA oligo of this invention comprises a desadenosine oligonucleotide, whether an anti-sense to a naturally occurring desthymidine sequence, or by substitution with one or more universal bases in accordance with the invention. The methods for substituting nucleotide, as well as for synthesizing

oligonucleotides of a specific sequence, and which bases to employ as universal bases are known in the art, and need not be further provided here, since they are within the knowledge

of an artisan.

In a further embodiment of the agent of the invention, the MTA oligo is operatively linked to an agent or molecule which, itself, is internalized or up-taken by living cells. In this manner, the uptake of the agent of the invention is enhanced as is known in the art. Examples of agents or molecules suitable for use with the MTA oligos of this invention are transferrin, asialoglycoprotein, and streptavidin. Others, however, are also suitable.

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The present agents are also provided as a pharmaceutical composition comprising an anti-sense oligonucleotide as given above in an amount effective to reduce expression of a target mRNA, by passing through a cell membrane and binding specifically with target mRNA in the cell so as to prevent its translation are another aspect of the present invention. Such compositions are provided in a suitable pharmaceutically acceptable carrier, e.g. sterile pyrogen-free saline solution. The agent of the invention may be formulated with a hydrophobic carrier capable of passing through a cell membrane, e.g. in a liposome, with the liposomes

carried in a pharmaceutically acceptable aqueous carrier. The oligonucleotides may be coupled to an agent which inactivates mRNA, such as a ribozyme. Such oligonucleotides may be administered to a subject in need of such treatment to inhibit the activation of specific receptors, enzymes and/or proteins and/or factors, among other expression products. The pharmaceutical formulation may also comprise chimeric molecules comprising anti-sense oligonucleotides attached to molecules which are known to be internalized by cells. These oligonucleotide conjugates utilize cellular up-take pathways to increase intracellular concentrations of the oligonucleotide. Examples of molecules used in this manner are macromolecules including transferrin, asialoglycoprotein (bound to oligonucleotides via polylysine) and streptavidin, among others.

The anti-sense compound may be contained in the pharmaceutical formulation within a lipid particle or vesicle, such as a liposome or microcrystal. The particles may be of any suitable structure, such as unilamellar or plurilamellar. The one preferred embodiment, the anti-sense oligonucleotide is comprised within the liposome. Positively charged lipids such as N-[1-(2, 3 -dioleoyloxi) propyl] -N, N, N-trimethylammoniumethylsulfate, or "DOTAP," are particularly preferred for such particles and vesicles. However, others are also suitable. The preparation of such lipid particles is well known. See, e.g., US Patent Nos. 4,880,635 to Janoff et al., 4,906,477 to Kurono et al., 4,911,928 to Wallach, 4,917,951 to Wallach, 4,920,016 to Allen et al., 4,921,757 to Wheatley et al., the relevant sections of all of which are herein incorporated in their entireties by reference.

The composition of the invention may be administered by any means which transports the agent to the lungs. The present agents may be administered to the lungs of a patient by any suitable means, but are preferably administered through the respiratory system as a respirable formulation, more preferably in the form of an aerosol comprising respirable particles which, in turn, comprise the agent for respiration or inhalation by the subject. The respirable particles may be in gaseous, liquid or solid form, and they may, optionally, contain other therapeutic ingredients and formulation components.

The particles of the present invention are preferably particles of respirable size, preferably of a size sufficiently small to pass, upon inhalation, through the mouth and larynx and into the bronchi and alveoli of the lungs. In general, particles ranging from about 0.5 to 10 microns in diameter are respirable. However, other sizes may also be suitable. Particles of non-respirable size, of considerably larger diameter, which are included in the respirable formulation tend to deposit in the throat and may be swallowed. Accordingly, it is desirable to minimize the quantity of non-respirable particles in the aerosol. For nasal administration, a particle size in the range of 10-500 μ m is preferred to ensure their retention in the nasal cavity.

Liquid pharmaceutical compositions of the agent of the invention for producing a respirable formulation, e.g. an aerosol may be prepared by combining the anti-sense oligo with a suitable vehicle or carrier, such as sterile pyrogen-free water and/or other known pharmaceutically or veterinarily acceptable carrier. Other therapeutic compounds may be included as well as other formulation ingredients as is known in the art.

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Solid particulate compositions comprising respirable dry particles of, e.g. the micronized agent of the invention may be prepared by grinding the dry anti-sense compound with a mortar and pestle, and then passing the thus ground, e.g. micronized composition through a screen, e.g. 400 mesh screen, to break up or separate large agglomerates of particles. A solid particulate composition comprising the anti-sense compound may optionally also comprise a dispersant and other known agents, which serve to facilitate the formation of a mist or aerosol. A suitable dispersant is lactose, which may be blended with the anti-sense compound in any suitable ratio, about 1:1 w/w. Other ratios may be utilized as well, and other therapeutic and formulation agents may also be included.

Aerosols of liquid particles comprising the agent may be produced by any suitable means, such as with an insufflator or nebulizer. See, e.g., US Patent No. 4,501,729. Nebulizers are commercially available devices which transform solutions or suspensions of an agent into a therapeutic aerosol mist either by means of acceleration of a compressed gas, typically air or oxygen, e.g. through a narrow venturi orifice or by means of ultrasonic agitation. Suitable formulations for use in insufflators and nebulizers comprise the present agent, the agent of this invention, in an amount of about 0.01 to about 40%, preferably less than 20% w/w in a liquid carrier which is typically water or a dilute aqueous alcoholic solution, preferably made isotonic with body fluids by the addition of, for example, sodium chloride. Other carriers are also suitable. Optional additives include preservatives if the formulation is not prepared sterile, for example, methyl hydroxybenzoate, antioxidants, flavoring agents, volatile oils, buffering agents and surfactants, among others.

The pharmaceutical compositions provided herein comprise nucleic acid(s) comprising the anti-sense oligonucleotide(s) described above and one or more surfactants. Suitable surfactants or surfactant components for enhancing the uptake of the anti-sense oligonucleotides of the invention include synthetic and natural as well as full and truncated forms of surfactant protein A, surfactant protein B, surfactant protein C, surfactant protein D and surfactant dipalmitoyl), phosphatidylcholine (other than Protein E. di-saturated phosphatidylcholine, dipalmitoylphosphatidylcholine, phosphatidylglycerol, phosphatidylinositol, phosphatidylethanolamine, phosphatidylserine; phosphatidic acid, lysophosphatidylethanolamine, lysophosphatidylcholine, ubiquinones, palmitoyl-lysophosphatidylcholine, dehydroepiandrosterone, dolichols, sulfatidic acid,

glycerol-3-phosphate, dihydroxyacetone phosphate, glycerol, glycero-3-phosphocholine, dihydroxyacetone, palmitate, cytidine diphosphate (CDP) diacylglycerol, CDP choline, choline, choline phosphate; as well as natural and artificial lamelar bodies which are the natural carrier vehicles for the components of surfactant, omega-3 fatty acids, polyenic acid, polyenoic acid, lecithin, palmitinic acid, non-ionic block copolymers of ethylene or propylene oxides, polyoxypropylene, monomeric and polymeric, polyoxyethylene, monomeric and polymeric, poly (vinyl amine) with dextran and/or alkanoyl side chains, Brij 35, Triton X-100 and synthetic surfactants ALEC, Exosurf, Survan and Atovaquone, among others. These surfactants may be useed either as single or part of a multiple component surfactant in a formulation, or as covalently bound additions to the 5' and/or 3' ends of the anti-sense oligonucleotides (oligos).

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The composition of the invention may be administered by any means which transports the anti-sense nucleotide and the surfactant composition to the lung. The anti-sense compounds disclosed herein may be administered to the lungs of a patient by any suitable means, but are preferably administered by inhalation of an aerosol comprised of respirable particles which comprise the anti-sense compound. The respirable particles may be liquid or solid, and they may optionally contain other therapeutic or diagnostic ingredients as well as other typical ingredients for a particular formulation. Examples of other agents are analgesics such as acetaminophen, anilerdine, aspirin, buprenorphine, butabital, butorpphanol, Choline Salicylate, Codeine, Dezocine, Diclofenac, Diflunisal, Dihydrocodeine, Elcatoninin, Etodolac, Fenoprofen, Hydrocodone, Hydromorphone, Ibuprofen, Ketoprofen, Ketorolac, Levorphanol, Magnesium Salicylate, Meclofenamate, Mefenamic Acid, Meperidine, Methadone, Methotrimeprazine, Morphine, Nalbuphine, Naproxen, Opium, Oxycodone, Oxymorphone, Pentazocine, Phenobarbital, Propoxyphene, Salsalate, Sodium Salicylate, Tramadol and Narcotic analgesics in addition to those listed above. See, Mosby's Physician's GenRx. Antianxiety agents are also useful including Alprazolam, Bromazepam, Buspirone, Chlordiazepoxide, Chlormezanone, Clorazepate, Diazepam, Halazepam, Hydroxyzine, Ketaszolam, Lorazepam, Meprobamate, Oxazepam and Prazepam, among others. Anti-anxiety agents associated with mental depression, such as Chlordiazepoxide, Amitriptyline, Loxapine Maprotiline and Perphenazine, among others. Anti-inflammatory agents such as non-rheumatic Aspirin, Choline Salicylate, Diclofenac, Diflunisal, Etodolac, Fenoprofen, Floctafenine, Flurbiprofen, Ibuprofen, Indomethacin, Ketoprofen, Magnesium Salicylate, Meclofenamate, Mefenamic Acid, Nabumetone, Naproxen, Oxaprozin, Phenylbutazone, Piroxicam, Salsalate, Sodium Salicylate, Sulindac, Tenoxicam, Tiaprofenic Acid, Tolmetin, anti-inflammatories for ocular treatment such as Diclofenac, Flurbiprofen, Indomethacin, Ketorolac, Rimexolone (generally for post-operative treatment), anti-inflammatories for, non-infectious nasal

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such as Beclomethaxone, Budesonide, Dexamethasone. Flunisolide. applications Triamcinolone, and the like. Soporifics (anti-insomnia/sleep inducing agents) such as those utilized for treatment of insomnia, including Alprazolam, Bromazepam, Diazepam, Diphenhydramine, Doxylamine, Estazolam, Flurazepam, Halazepam, Ketazolam, Lorazepam, Nitrazepam, Prazepam Quazepam, Temazepam, Triazolam, Zolpidem and Sopiclone, among including Diphenhydramine, Hydroxyzine, Sedatives Promethazine, Propofol, Melatonin, Trimeprazine, and the like. Sedatives and agents used for treatment of petit mal and tremors, among other conditions, such as Amitriptyline HCl; Chlordiazepoxide, Amobarbital; Secobarbital, Aprobarbital, Butabarbital, Ethchiorvynol, Glutethimide, L-Tryptophan, Mephobarbital, MethoHexital Na, Midazolam Hcl, Oxazepam, Pentobarbital Na, Phenobarbital, Secobarbital Na, Thiamylal Na, and many others. Agents used in the treatment of head trauma (Brain Injury/Ischemia), such as Enadoline HCl (e.g. for treatment of severe head injury; orphan status, Warner Lambert), cytoprotective agents, and agents for the treatment of menopause, menopausal symptoms (treatment), e.g. Ergotamine, Belladonna Alkaloids and Phenobarbital, for the treatment of menopausal vasomotor symptoms, e.g. Clonidine, Conjugated Estrogens and Medroxyprogesterone, Estradiol, Estradiol Cypionate, Estradiol Valerate, Estrogens, conjugated Estrogens, esterified Estrone, Estropipate, and Ethinyl Estradiol. Examples of agents for treatment of pre menstrual syndrome (PMS) are Progesterone, Progestin, Gonadotrophic Releasing Hormone, Oral contraceptives, Danazol, Luprolide Acetate,

Vitamin B6. Examples of agents for treatment of emotional/psychiatric treatments such as Tricyclic Antidepressants, including Amitriptyline HCl (Elavil), Amitriptyline HCl, Perphenazine (Triavil) and Doxepin HCl (Sinequan). Examples of tranquilizers, antidepressants and anti-anxiety agents are Diazepam (Valium), Lorazepam (Ativan), Alprazolam (Xanax), SSRI's (selective Ssrotonin reuptake inhibitors), Fluoxetine HCl (Prozac), Sertaline HCl (Zoloft), Paroxetine HCl (Paxil), Fluoxamine Maleate (Luvox), Venlafaxine HCl (Effexor), Serotonin, Serotonin Agonists (Fenfluramine), and other over the counter (OTC) medications.

The composition of the present invention may be administered into the respiratory system as a formulation including particles of respirable size, e.g. particles of a size sufficiently small to pass through the nose, mouth and larynx upon inhalation and through the bronchi and alveoli of the lungs. In general, respirable particles range from about 0.5 to 10 microns in size. Particles of non-respirable size which are included in the aerosol tend to deposit in the throat and be swallowed, and the quantity of non-respirable particles in the aerosol is thus minimized. For nasal administration, a particle size in the range of 10-500 μ m is preferred to ensure retention in the nasal cavity.

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Aerosols or mists of solid particles comprising the agent of the invention may likewise be produced with any device that generates solid particulate medicament aerosols or mists. Aerosol and mist generators are suitable for administering solid particulate medicaments. These devices produce respirable particles, as explained above., and generate a volume of aerosol or mist containing a predetermined metered dose of a medicament at a rate suitable for human or animal administration. One illustrative type of solid particulate aerosol generator is an insufflator. Suitable formulations for administration by insufflation include finely comminuted powders which may be delivered by means of an insufflator or taken into the nasal cavity in the manner of a snuff. In the insufflator, the powder, e.g. a metered dose of the agent effective to carry out the treatments described herein, is contained in a capsule or a cartridge. These capsules or cartridges are typically made of gelatin or plastic, and may be pierced or opened in situ, and the powder delivered by air drawn through the device upon inhalation or by means of a manually-operated pump. The powder employed in the insufflator may consist either solely of the agent or of a powder blend comprising the agent, a suitable powder diluent, such as lactose, and an optional surfactant as well as other agents. The agent typically comprises from 0.01 to 100 w/w of the formulation. A second type of illustrative aerosol generator comprises a metered dose inhaler. Metered dose inhalers are pressurized aerosol dispensers, typically comprising a suspension or solution formulation of the active ingredient in a liquified propellant. During use these devices discharge the formulation through a valve adapted to deliver a metered volume, typically about 10 to 150 μ l, although other volumes are also suitable, to produce a fine particle spray containing the active ingredient. Suitable propellants include solvents such as certain chlorofluorocarbon compounds, for example, dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane and/or mixtures thereof. The formulation may additionally comprise one or more co-solvents, for example, ethanol, surfactants, such as oleic acid or sorbitan trioleate, antioxidants and suitable flavoring agents. The aerosol, whether formed from solid or liquid particles, may be produced by the aerosol generator at a rate of from about 10 to 150 liters per minute, more preferably from about 30 to 150 liters per minute, and most preferably about 60 liters per minute. Aerosols containing greater amounts of medicament may be administered more rapidly.

As already indicated, the agent of this invention is also provided as a composition, comprising the agent of the invention, and a carrier. The carrier is preferably a biologically acceptable carrier, and more preferably a pharmaceutically or veterinarily acceptable carrier in the form of a gaseous, liquid, solid carriers, and mixtures thereof, which are suitable for the different routes of administration intended. The composition may optionally comprise other agents such as other therapeutic compounds known in the art for the treatment of the condition or disease, antioxidants, flavoring and coloring agents, fillers, volatile oils, buffering agents,

dispersants, surfactants, RNA inactivating agents, antioxidants, flavoring agents, propellants and preservatives, as well as other agents known to be utilized in therapeutic compositions. An example of the mRNA inactivating agent is an enzyme, such as ribozyme.

The composition generally contains the anti-sense oligonucleotide in an amount of about 0.01 to about 99.99 w/w, preferably about 1 to about 40 w/w, and more preferably about 5 to about 20 w/w of the composition. However, other ingredients, and other amounts of the agent are also suitable within the confines of this invention.

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The agent of the invention is also provided in various formulations which are tailored for different methods of administration and routes of delivery. The formulations that are contemplated are, for example, a transdermal formulation also containing carrier(s) and other agents suitable for delivery through the skin, mouth, nose, vagina, anus, eyes, ears, other body cavities, intradermally, as a sustained release formulation, intracranial, intrathecally, intravascularly, by inhalation, intrapulmonarily, into an organ, by implantation, including suppositories, cremes, gels, and the like, as is known in the art. In one particular formulation, the agent is suspended or dissolved in a solvent. In another the carrier comprises a hydrophobic carrier, such as lipid particles or vesicles, including liposomes and micro crystals. The preparation of all of these formulations, as well as the ingredients to be utilized are known in the art, and need not be further described here. In one particularly preferred embodiment of the vesicles comprise liposomes containing the anti-sense the vesicle formulation, oligonucleotide. The lipid vesicles may comprise N-(1-[2, 3-dioleoxyloxi] propyl) -N,N,Ntrimethyl- ammonium methylsulfate as well as other lipids known in the art to provide suitable delivery of DNA to target cells. In one embodiment, the formulation comprises a respirable formulation, such as an aerosol. The agent, composition, and formulation of the invention are provided in bulk, and in unit form, as well as in the form of an implant, a solution or suspension, a capsule or cartridge, which may be openable or piercable as is known in the art.

A kit is also provided, which comprises a delivery device, and in separate containers, the agent, composition or formulation of the invention, and optionally other agents, and instructions for the use of the kit components. In one preferred embodiment, the delivery device comprises a nebulizer which delivers single or multiple metered doses of the formulation. The single metered dose nebulizer may be provided as a disposable kit which is sterilely preloaded with enough agent for one application. The nebulizer may be provided as an insufflator, and the composition in a piercable or openable capsule or cartridge. In a different embodiment, the delivery device comprises a pressurized inhaler, and the agent is in the form of a suspension or solution. The kit may optionally also comprise in a separate container an agent selected from the group consisting of other therapeutic compounds, antioxidants, flavoring and coloring agents, fillers, volatile oils, buffering agents, dispersants,

surfactants, cell internalized or up taken agents, RNA inactivating agents, antioxidants, flavoring agents, propellants and preservatives, among other suitable additives for the different formulations. When a solvent for the agent or the other ingredients is added, organic solvents and organic solvents mixed with one or more co-solvents may be utilized as well as aqueous solvents as is known in the art.

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The agent of the invention may be provided in conjunction with a vector for delivery purposes, or for manufacturing copies thereof. The agent may be operatively linked to the vector as is known in the art. The agent may also be provided within a host cell for amplification of the MTA oligo, and for storage purposes.

The agent of this invention may be utilized by itself or in the form of a composition or various formulations in the treatment of a disease or condition associated with the mRNA corresponding to at least one target gene(s), to genomic flanking regions, initiation codon, intron-exon borders and the like, or the entire sequence of precursor RNAs, including noncoding RNA segments, the 5'-end and the 3'-end, e.g. poly-A segment and oligos targeted to the juxta-section between coding and non-coding regions, and RNA regions encoding proteins, by administration to a subject afflicted with the disease or condition of an amount of the antisense oligonucleotide effective to reduce the production or availability, or to increase the degradation by the subject of at least one of the target mRNAs. Typically, the agent is administered in an amount effective to reduce the production or availability, or to increase the degradation of at least two of the target mRNAs. Optionally, the agent is administered directly to the lung (s) of the subject, preferably as a respirable aerosol. Although an artisan will know how to titrate the amount of agent to be administered by the weight of the subjected being treated in accordance with the teachings of this patent, the agent is preferably administered in an amount effective to attain an intracellular concentration of about 0.05 to about 10 μM multiple targeted anti-sense oligonucleotide, preferably in an amount effective to attain an intracellular concentration of up to about 5 μM multiple targeted anti-sense oligonucleotide.

The treatment provided in this patent is suitable for treating numerous diseases and conditions, and its application is solely limited by the availability of target molecules and their sequences. One type of disease or condition for which this technology is particularly well suited are lung diseases or conditions. For this type of application, at least one of the target mRNA encodes a protein such as the adenosine A₁ receptor, adenosine A₂B receptor, adenosine A₃ receptor, and bradykinin B2 receptor. However, other targets are also suitable. In one application, the disease or condition is associated with obstruction of the subject's airways, in another specifically with asthma, etc. One of the preferred target proteins comprises the NfkB transcription factor, although others which were described above are also suitable. In another preferred application, the disease or condition is associated with inflammation. For this type

of application at least one of the target mRNA preferably encodes a protein selected from the group consisting of interleukins, chemokines, Rantes and receptors thereof. Still another application is for treating a disease or condition associated with an allergy. For this application, the mRNA preferably encodes a target selected from the group consisting of an antibody and an antibody receptor. For the application of this technology to a disease or condition associated with a malignancy or cancer, the mRNA preferably encodes a target selected from the group consisting of oncogenes, an immunoglobulin and an antibody receptor.

Depending on the target organ or tissue, the agent of the invention may be delivered in one of many ways, for example, by a transdermal or systemic route, and more specifically orally, intracavitarily, intranasally, intraanally, intravaginally, transdermally, intrabucally, intravenously, subcutaneously, intramuscularly, intratumorously, into a gland, by implantation, intradermally, and many other routes of administration. The formulation may be, in addition, an implant, slow release, transdermal release, sustained release, and coated with one or more macromolecules to avoid destruction of the agent prior to reaching the selected target. The subject that may be treated by the present agent are varied, and include humans and other animals in general, and in particular vertebrates, and amongst these mammals, and more specifically humans, and small and large, wild and domesticated, marine and farm animals, and preferably humans and domesticated and farm animals. In one aspect of the invention, at least one of the target mRNAs and the subject are of the same species, and in a preferred case they are of human origin. However, since in one embodiment mismatched nucleotides are replaced, mismatched species may also be utilized.

The multiple targeted anti-sense oligonucleotide of this invention may be administered in a broad dose range. Preferable is an amount of about 0.005 to about 150 mg/kg body weight per administration, and the agent may be administered from once in an acute treatment to several doses per day, to a continuous administration to maintain the level of a specific molecule. Preferred doses are about 0.01 to about 75 mg/kg body weight, more preferably about 1 to 50 mg/kg body weight. The method may be administered as a prophylactic or therapeutic method.

The agent of the invention may be produced by selecting two or more targets selected from the group consisting of genes, genomic flanking regions, mRNAs and proteins known to be associated with at least one disease or condition; obtaining RNAs selected from the group consisting of RNAs corresponding to the genes, to genomic flanking regions, initiation codon, intron-exon borders and the like, or the entire sequence of RNAs, including non-coding RNA segments, the 5'-end and the 3'-end, e.g. the poly-A segment and oligos targeted to the juxta-section between coding and non-coding regions, and RNA segments encoding the target proteins; selecting a segment of a first RNA which is at least about 60% homologous to a

segment of at least a segment of a second RNA; and synthesizing one or more anti-sense oligonucleotide(s) to the one or more RNA segments. In one preferred embodiment, the method further comprises substituting a universal base for at least one, and in some instances all, non-homologous nucleotide in the anti-sense oligonucleotide, and in another preferred embodiment the method further comprises substituting a methylated cytosine for cytosine in at least one CpG dinucleotide present in the anti-sense oligonucleotide. The technology involved in methylation is known in the art and need not be further described here.

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Although the specific length of the MTA oligo is determined by the target's length, and its segments containing few thymidines, the anti-sense oligonucleotide(s) are preferably greater than about 7 nucleotides long, and up to about 60 nucleotides long, and longer. The specific backbone chemistry may by selected by an artisan based on the teachings provided here and the knowledge of the art at large. One factor that impinges on the selection of the nucleotide bridging residues is the level of nuclease resistance desired and other factors specific to one or the other method of administration. Another factor is the need for localization of the treatment, to minimize or fully avoid side effects which might otherwise be caused along with the therapeutic effect of the agent.

The following examples are provided to illustrate the present invention, and should not be construed as limiting thereon. In these examples, μ M means micromolar, ml means milliliters, μ M means micrometers, mm means millimeters, cm means centimeters, °C means degrees Celsius, μ g means micrograms, mg means milligrams, g means grams, kg means kilograms, M means molar, and h means hours.

EXAMPLES

Example 1: Design and Synthesis of Anti-sense Oligonucleotides

The design of anti-sense oligonucleotides against the A₁ and A₂ adenosine receptors may require the solution of the complex secondary structure of the target A₁ receptor mRNA and the target A₃ receptor mRNA. After generating this structure, anti-sense nucleotide are designed which target regions of mRNA which might be construed to confer functional activity or stability to the mRNA and which optimally may overlap the initiation codon. Other target sites are readily usable. As a demonstration of specificity of the anti-sense effect, other oligonucleotides not totally complementary to the target mRNA, but containing identical nucleotide compositions on a w/w basis, are included as controls in anti-sense experiments.

The mRNA secondary structure of the adenosine A₁ receptor was analyzed and used as described above, to design a phosphorothioate anti-sense oligonucleotide. The anti-sense oligonucleotide which was synthesized was designated HAdA_iAS and had the following sequence:

5' -GAT GGA GGG CGG CAT GGC GGG-3' (SEQ ID NO:1)

As a control, a mismatched phosphorothioate anti-sense nucleotide designated HAdAlMM1 was synthesized with the following sequence:

5' -GTA GCA GGC GGG GAT GGG GGC-3' (SEQ ID NO:2)

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Each oligonucleotide had identical base content and general sequence structure. Homology searches in GENBANK (release 85.0) and EMBL (release 40.0) indicated that the anti-sense oligonucleotide was specific for the human and rabbit adenosine A_i receptor genes, and that the mismatched control was not a candidate for hybridization with any known gene sequence.

The secondary structure of the adenosine A₃ receptor mRNA was similarly analyzed and used as described above to design two phosphorothioate anti-sense oligonucleotides. The first anti-sense oligonucleotide (HAdA3AS1) synthesized had the following sequence:

5' -GTT GTT GGG CAT CTT GCC-3' (SEQ ID NO:3)

As a control, a mismatched phosphorothioate anti-sense oligonucleotide (HAdA3MM1) was synthesized, having the following sequence:

5' -GTA CTT GCG GAT CTA GGC-3' (SEQ ID NO:4)

A second phosphorothioate anti-sense oligonucleotide (HAdA3AS2) was also designed and synthesized, having the following sequence:

5' -GTG GGC CTA GCT CTC GCC-3' (SEQ ID NO:5)

Its control oligonucleotide (HAdA3MM2) had the sequence:

5' -GTC GGG GTA CCT GTC GGC-3' (SEQ ID NO:6)

Phosphorothioate oligonucleotides were synthesized on an Applied Biosystems Model 396 Oligonucleotide Synthesizer, and purified using NENSORB chromatography (DuPont, MD).

25 Example 2: In Vivo Testing of Adenosine A₁ Receptor Anti-sense Oligos

The anti-sense oligonucleotide against the human A₁ receptor (SEQ ID NO:1) described above, was tested for efficacy in an in vitro model utilizing lung adenocarcinoma cells HTB-54, HTB-54 lung adenocarcinoma cells were demonstrated to express the A₁ adenosine receptor using standard northern blotting procedures and receptor probes designed and synthesized in the laboratory.

HTB-54 human lung adenocarcinoma cells (106/100 mm tissue culture dish) were exposed to $5.0 \,\mu\text{M}$ HAdAlAS or HAdAlMM1 for 24 hours, with a fresh change of media and oligonucleotides after 12 hours of incubation. Following 24 hour exposure to the oligonucleotides, cells were harvested and their RNA extracted by standard procedures. A 21-mer probe corresponding to the region of mRNA targeted by the anti-sense (and therefore

having the same sequence as the anti-sense, but not phosphorothioated) was synthesized and used to probe northern blots of RNA prepared from HAdAlAS-treated, HAdAlMM1-treated and non-treated HTB-54 cells. These blots showed clearly that HAdAlAS but not HAdAlMM1 effectively reduced human adenosine receptor mRNA by >50%. This result showed that HAdAlAS is a good candidate for an anti-asthma drug since it depletes intracellular mRNA for the adenosine A_1 receptor, which is involved in asthma.

Example 3: In Vivo Efficacy of Adenosine A₁ Receptor Anti-sense Oligos

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A fortuitous homology between the rabbit and human DNA sequences within the adenosine A_1 gene overlapping the initiation codon permitted the use of the phosphorothioate anti-sense oligonucleotides initially designed for use against the human adenosine A_1 receptor in a rabbit model.

Neonatal New Zealand white Pasteurella-free rabbits were immunized intraperitoneally within 24 hours of birth with 312 antigen units/ml house dustmite (D. farinae) extract (Berkeley Biologicals, Berkeley, CA), mixed with 10% kaolin. Immunizations were repeated weekly for the first month and then biweekly for the next 2 months. At 3-4 months of age, eight sensitized rabbits were anesthetized and relaxed with a mixture of ketamine hydrochloride (44 mg/kg) and acepromazine maleate (0.4 mg/kg) administered intramuscularly.

The rabbits were then laid supine in a comfortable position on a small molded, padded animal board and intubated with a 4.0-mm intratracheal tube (Mallinkrodt, Inc., Glens Falls, NY). A polyethylene catheter of external diameter 2.4 mm with an attached latex balloon was passed into the esophagus and maintained at the same distance (approximately 16 cm) from the mouth throughout the experiments. The intratracheal tube was attached to a heated Fleisch pneumotachograph (size 00; DOM Medical, Richmond, VA), and flow was measured using a Validyne differential pressure transducer (Model DP-45161927; Validyne Engineering Corp., Northridge, CA) driven by a Gould carrier amplifier (Model 11-4113; Gould Electronic, Cleveland, OH). The esophageal balloon was attached to one side of the differential pressure transducer, and the outflow of the intratracheal tube was connected to the opposite side of the pressure transducer to allow recording of transpulmonary pressure. Flow was integrated to give a continuous tidal volume, and measurements of total lung resistance (RL) and dynamic compliance (Cdyn) were calculated at isovolumetric and flow zero points, respectively, using an automated respiratory analyzer (Model 6; Buxco, Sharon, CT).

Animals were randomized and on Day 1 pretreatment values for PC50 were obtained for aerosolized adenosine. Anti-sense (HAdAlAS) or mismatched control (HAdAlMM) oligonucleotides were dissolved in sterile physiological saline at a concentration of 5000 μ g

(5 mg) per 1.0 ml. Animals were subsequently administered the aerosolized anti-sense or mismatch oligonucleotide via the intratracheal tube (approximately 5000 μ g in a volume of 1.0 ml), twice daily for two days. Aerosols of either saline, adenosine, or anti-sense or mismatch oligonucleotides were generated by an ultrasonic nebulizer (DeVilbiss, Somerset, PA), producing aerosol droplets 80% of which were smaller than 5 μ m in diameter.

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In the first arm of the experiment, four randomly selected allergic rabbits were administered anti-sense oligonucleotide and four the mismatched control oligonucleotide. On the morning of the third day, PC50 values (the concentration of aerosolized adenosine in mg/ml required to reduce the dynamic compliance of the bronchial airway 50% from the baseline value) were obtained and compared to PC50 values obtained for these animals prior to exposure to oligonucleotide.

Following a 1 week interval, animals were crossed over, with those previously administered mismatch control oligonucleotide now administered anti-sense oligonucleotide, and those previously treated with anti-sense oligonucleotide now administered mismatch control oligonucleotide. Treatment methods and measurements were identical to those employed in the first arm of the experiment. It should be noted that in six of the eight animals treated with anti-sense oligonucleotide, adenosine-mediated bronchoconstriction could not be obtained up to the limit of solubility of adenosine, 20 mg/ml. For the purpose of calculation, PC50 values for these animals were set at 20 mg/ml. The values given therefore represent a minimum figure for anti-sense effectiveness. Actual effectiveness was higher. The results of this experiment are illustrated in Table 3 below.

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Effect of Adenosine A, Receptor Anti-sense Oligo Table 3: upon PC50 Values in Asthmatic Rabbits

Misma	tch Control	A ₁ Receptor An	A ₁ Receptor Anti-sense Oligo		
Pre Oligonucleotide	Post Oligonucleo	otide Pre Oligonucleotide	Post Oligonucleotide		
	5.16 ± 1.03	2.36 ± 0.68	>19.5 ± 0.34**		

The results are presented as the mean $(n=8) \pm SEM$.

The significance was determined by repeated-measures analysis of variance (ANOVA), and Tukey's protected test.

**Significantly different from all other groups, p<0.01.

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In both arms of the experiment, animals receiving the anti-sense oligonucleotide showed an order of magnitude increase in the dose of aerosolized adenosine required to reduce dynamic compliance of the lung by 50%. No effect of the mismatched control oligonucleotide upon PC50 values was observed. No toxicity was observed in any animal receiving either anti-sense or control inhaled oligonucleotide.

These results show clearly that the lung has exceptional potential as a target for antisense oligonucleotide-based therapeutic intervention in lung disease. They further show, in a model system which closely resembles human asthma, that downregulation of the adenosine A₁ receptor largely eliminates adenosine-mediated bronchoconstriction in asthmatic airways. Bronchial hyperresponsiveness in the allergic rabbit model of human asthma is an excellent endpoint for anti-sense intervention since the tissues involved in this response lie near to the point of contact with aerosolized oligonucleotides, and the model closely simulates an important human disease.

Specificity of A1-adenosine Receptor Anti-sense Oligonucleotide

At the conclusion of the cross-over experiment of Example 3 above, airway smooth muscle from all rabbits was quantitatively analyzed for adenosine A₁ receptor number. As a control for the specificity of the anti-sense oligonucleotide, adenosine A2 receptors, which should not have been affected, were also quantified.

Airway smooth muscle tissue was dissected from each rabbit and a membrane fraction prepared according to the method of Kleinstein et al. (Kleinstein, J. and Glossmann, H., Naunyn-Schmiedeberg's Arch. Pharmacol. 305: 191-200 (1978)), the relevant portion of which is hereby incorporated in its entirety by reference, with slight modifications. Crude plasma membrane preparations were stored at 70°C until the time of assay. Protein content was determined by the method of Bradford (M. Bradford, Anal. Biochem. 72, 240-254 (1976), the relevant portion of which is hereby incorporated in its entirety by reference). Frozen plasma

membranes were thawed at room temperature and were incubated with 0.2 U/ml adenosine deaminase for 30 minutes at 37°C to remove endogenous adenosine. The binding of [³H] DPCPX (A₁ receptor-specific) or [³H] CGS-21680 (A₁ receptor-specific) was measured as previously described by Ali et al. (Ali, S. et al., J. Pharmacol. Exp. Ther. 268, Am. J. Physiol 266, L271-277 (1994), the relevant portion of which is hereby incorporated in its entirety by reference).

The animals treated with adenosine A_1 anti-sense oligonucleotide in the cross-over experiment had a nearly 75% decrease in A_1 receptor number compared to controls, as assayed by specific binding of the A_1 -specific antagonist DPCPX. There was no change in adenosine A_2 receptor number, as assayed by specific binding of the A_2 receptor-specific agonist 2- [p-(2-carboxyethyl)-phenethylamino] -5' - (N-ethylcarboxamido) adenosine (CGS-21680). This is illustrated in Table 4 below.

Table 4: Specificity of Action of Adenosine A, Receptor Anti-sense Oligonucleotide

	Mismatch Control Oligonucleotide	A ₁ Anti-sense Oligonucleotide	
A ₁ -Specific Binding	1105 ± 48**	293 ± 18	
A ₂ -Specific Binding	302 ± 22	442 ± 171	

The results are presented as the mean $(n = 8) \pm SEM$.

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The significance was determined by repeated-measures analysis of variance (ANOVA), and Tukey's protected test.

**Significantly different from mismatch control, p<0.01.

The above results illustrate the effectiveness of anti-sense oligonucleotides in treating airway disease. Since the anti-sense oligos described above, eliminate the receptor systems responsible for adenosine-mediated bronchoconstriction, it may be less imperative to eliminate adenosine from them. However, it would be preferable to eliminate adenosine from even these oligonucleotides to reduce the dose needed to attain a similar effect. Described above are other anti-sense oligonucleotides targeting mRNA of proteins involved in inflammation. Adenosine has been eliminated from their nucleotide content to prevent its liberation during degradation.

Example 5: Anti-sense Oligos directed to other Target Nucleic Acids

This work was conducted to demonstrate that the present invention is broadly applicable to anti-sense oligonucleotides ("oligos") specific to nucleic acid targets broadly. The following experimental studies were conducted to show that the method of the invention is broadly suitable for use with anti-sense oligos designed as taught by this application and targeted to any

and all adenosine receptor mRNAs. For this purpose, various anti-sense oligos were porepared to adenosine receptor mRNAs exemplified by the adenosine A_1 , A_{2b} and A_3 receptor mRNAs.

Anti-sense Oligo I was disclosed above (SEQ. ID NO:1). Five additional anti-sense phosphorothicate oligos were designed asnd synthesized as indicated above.

- 1- Oligo II (SEQ. ID NO: 7) also targeted to the adenosine A₁ receptor, but to a different region than Oligo I.
- 2-Oligo V (SEQ. ID NO: 10) targeted to the adenosine A_{2b} receptor.
- 3- Oligos III (SEQ. ID NO: 8) and IV (SEQ. ID NO: 9) targeted to different regions of the adenosine A_3 receptor.
- 4- Oligo I-PD (SEQ. ID NO: 1681)(a phosphodiester oligo of the same sequence as Oligo I).

These anti-sense oligos were designed for therapy on a selected species as described above and are generally specific for that species, unless the segment of the target mRNA of other species happens to contain a similar sequences. All anti-sense oligos were prepared as described below, and tested in vivo in a rabbit model for bronchoconstriction, inflammation and allergy, which have breathing difficulties and impeded lung airways, as is the case in ailments such as asthma, as described in the above-identified application.

Example 6: Design & Sequences of other Anti-sense Oligos

Six oligos and their effects in a rabbit model were studied and the results of these studies are reported and discussed below. Five of these oligos were selected for this study to complement the data on Oligo I (SEQ ID NO: 1) provided in Examples 1 to 4 above. This oligo is anti-sense to one region of the adenosine A₁ receptor mRNA.

The oligos tested are identified as anti-sense Oligos I (SEQ ID NO: 1) and II (SEQ. ID No: 7) targeted to a different region of the adenosine A₁ receptor mRNA, Oligo V (SEQ. ID No:8) targeted to the adenosine A_{2b} receptor mRNA, and anti-sense Oligos III and IV (SEQ. ID NOS: 9 and 10) targeted to two different regions of the adenosine A₃ receptor mRNA. The sixth oligo (Oligo I-PD) is a phosphodiester version of Oligo I (SEQ. ID NO:1). The design and synthesis of these anti-sense oligos was performed in accordance with Example 1 above.

(I) Anti-sense Oligo I

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The anti-sense oligonucleotide I referred to in Examples 1 to 4 above is targeted to the human A_1 adenosine receptor mRNA (EPI 2010). Anti-sense oligo I is 21 nucleotide long, overlaps the initiation codon, and has the following sequence.

5'- GAT GGA GGG CGG CAT GGC GGG -3' (:SEQ. ID No 1)

The oligo I was previously shown to abrogate the adenosine-induced bronchoconstriction in allergic rabbits, and to reduce allergen-induced airway obstruction and

bronchial hyperresponsiveness (BHR), as discussed above and shown by Nyce, J. W. & Metzger, W. J., Nature, 385:721 (1977), the relevant portions of which reference are incorporated in their entireties herein by reference.

(II) Anti-sense Oligo II

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A phosphorothicate anti-sense oligo (SEQ. ID NO:7) was designed in accordance with the invention to target the rabbit adenosine A_1 receptor mRNA region +936 to +956 relative to the initiation codon (start site). The anti-sense oligo II is 21 nucleotide long, and has the following sequence.

5'-CTC GTC GCC GTC GCC GGC GGG-3' (SEQ. ID NO:7)

(III) Anti-sense Oligo III

A phosphorothicate anti-sense oligo other than that provided in Example 1 above (SEQ. ID NO:8) was designed in accordance with the invention to target the anti-sense A_3 receptor mRNA region +3 to + 22 relative to the initiation codon start site. The anti-sense oligo III is 20 nucleotide long, and has the following sequence.

5'-GGG TGG TGC TAT TGT CGG GC-3' (SEQ. ID NO:8)

(IV) Anti-sense Oligo IV

Yet another phosphorothioate anti-sense oligo (SEQ. ID NO:9) was designed in accordance with the invention to target the adenosine A_3 receptor mRNA region + 386 to + 401 relative to the initiation codon (start site). The anti-sense oligo IV is 15 nucleotide long, and has the following sequence.

5'-GGC CCA GGG CCA GCC-3' (SEQ. ID NO:9)

(V) Anti-sense Oligo V

A phosphorothioate anti-sense oligo (SEQ. ID NO:10) was designed in accordance with the invention to target the adenosine A_{2b} receptor mRNA region -21 to -1 relative to the initiation codon (start site). The anti-sense oligonucleotide V is 21 nucleotide long, and has the following sequence.

5'-GGC CGG GCC AGC CGG GCC CGG-3' (SEQ. ID NO:10)

(VI) A, Mismatch Oligos

Two different mismatched oligonucleotides having the following sequences were used as controls for anti-sense oligo I (SEQ. ID NO: 1) described in Example 5 above.

A₁ MM2 5'-GTA GGT GGC GGG CAA GGC GGG-3' (SEQ. ID NO:1682) A₁ MM3 5'-GAT GGA GGC GGG CAT GGC GGG-3' (SEQ. ID NO:1683)

Anti-sense oligo I and the two mismatch anti-sense oligos had identical base content and general sequence structure. Homology searches in GENBANK (release 85.0) and EMBL (release 40.0) indicated that the anti-sense oligo I was specific, not only for the human, but also for the rabbit, adenosine A_i receptor genes, and that the mismatched controls were not candidates for hybridization with any known human or animal gene sequence.

(VII) Anti-sense Oligo A₁-PD (Oligo VI)

A phosphodiester anti-sense oligo (Oligo VI; SEQ. ID NO:1681) having the same nucleotide sequence as Oligo I was designed as disclosed in the above-identified application. Anti-sense oligo I-PD is 21 nucleotide long, overlaps the initiation codon, and has the following sequence.

5'- GAT GGA GGG CGG CAT GGC GGG -3' (SEQ. ID NO:1681)

III) Controls

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Each rabbit was administered 5.0 ml aerosolized sterilé saline following the same schedule as for the anti-sense oligos in (II), (III), and (IV) above.

10 Example 7: Synthesis of Anti-sense Oligos

Phosphorothioate anti-sense oligos having the sequences described in (a) above, were synthesized on an Applied Biosystems Model 396 Oligonucleotide Synthesizer, and purified using NENSORB chromatography (DuPont, DE). TETD (tetraethylthiuram disulfide) was used as the sulfurizing agent during the synthesis. Anti-sense oligonucleotide II (SEQ. ID NO:7), anti-sense oligonucleotide III (SEQ. ID NO: 8) and anti-sense oligonucleotide IV (SEQ. ID NO: 9) were each synthesized and purified in this manner.

Example 8: Preparation of Allergic Rabbits

Neonatal New Zealand white Pasturella-free rabbits were immunized intraperitoneally within 24 hours of birth with 0.5 ml of 312 antigen units/ml house dust mite (D. farinae) extract (Berkeley Biologicals, Berkeley, CA) mixed with 10% kaolin as previously described (Metzger, W. J., in Late Phase Allergic Reactions, Dorsch, W., Ed., CRC Handbook, pp. 347-362, CRC Press, Boca Raton (1990); Ali, S., Metzger, W. J. and Mustafa, S. J., Am. J. Resp. Crit. Care Med. 149: 908 (1994)), the relevant portions of which are incorporated in their entireties here by reference. Immunizations were repeated weekly for the first month and then biweekly until the age of 4 months. These rabbits preferentially produce allergen-specific IgE antibody, typically respond to aeroallergen challenge with both an early and late-phase asthmatic response, and show bronchial hyper responsiveness (BHR). Monthly intraperitoneal administration of allergen (312 units dust mite allergen, as above) continues to stimulate and maintain allergen-specific IgE antibody and BHR. At 4 months of age, sensitized rabbits were prepared for aerosol administration as described by Ali et al. (Ali, S., Metzger, W. J. and Mustafa, S. J., Am. J. Resp. Crit. Care Med. 149 (1994)), the relevant section being incorporated in its entirety here by reference.

DOSE-RESPONSE STUDIES

Example 9: Experimental Setup

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Aerosols of either adenosine (0-20 mg/ml), or anti-sense or one of two mismatch oligonucleotides (5 mg/ml) were separately prepared with an ultrasonic nebulizer (Model 646, DeVilbiss, Somerset, PA), which produced aerosol droplets, 80% of which were smaller than 5μ m in diameter. Equal volumes of the aerosols were administered directly to the lungs via an intratracheal tube.

The animals were randomized, and administered aerosolized adenosine. Day 1 pretreatment values for sensitivity to adenosine were calculated as the dose of adenosine causing a 50% loss of compliance (PC_{50} Adenosine). The animals were then administered either the aerosolized anti-sense or one of the mismatch anti-sense oligos via the intratracheal tube (5 mg/1.0 ml), for 2 minutes, twice daily for 2 days (total dose, 20 mg). Post-treatment PC_{50} values were recorded (post-treatment challenge) on the morning of the third day. The results of these studies are provided in Example 21 below.

Example 10: Crossover Experiments

For some experiments utilizing anti-sense oligo I (SEQ ID NO: 1) and a corresponding mismatch control oligonucleotide A1MM2, following a 2 week interval, the animals were crossed over, with those previously administered the mismatch control A_1MM2 , now receiving the anti-sense oligo I, and those previously treated with the anti-sense oligo I, now receiving the mismatch control A_1MM2 oligo.

The number of animals per group was as follows. For mismatch A_iMM2 (Control 1), n=7, since one animal was lost in the second control arm of the experiment due to technical difficulties, for mismatch A_iMM3 n=4 (Control 2) and for A_iAS anti-sense oligo I, n=8. The A_iMM3 oligo-treated animals were analyzed separately and were not part of the cross-over experiment. The treatment methods and measurements employed following the cross-over were identical to those employed in the first arm of the experiment.

In 6 of the 8 animals treated with the anti-sense oligo I (SEQ. ID NO: 1), no PC₅₀ value could be obtained for adenosine doses of up to 20 mg/ml, which is the limit of solubility of adenosine. Accordingly, the PC₅₀ values for these animals were assumed to be 20 mg/ml for calculation purposes. The values given, therefore, represent a minimum figure for the effectiveness of the anti-sense oligonucleotides of the invention. Other groups of allergic rabbits (n=4 for each group) were administered 0.5 or 0.05 mg doses of the anti-sense oligo I (SEQ ID NO: 1), or the A₁MM2 oligo in the manner and according to the schedule described above (the total doses being 2.0 or 0.2 mg). The results of these studies are provided in Example 22 below.

Example 11: Anti-sense Oligo Formulation

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Each one of anti-sense oligos were separately solubilized in an aqueous solution and administered as described for anti-sense oligo I (SEQ. ID No:1) in (e) above, in four 5 mg aliquots (20 mg total dose) by means of a nebulizer via endotracheal tube, as described above.

The results obtained for anti-sense oligo I and its mismatch controls confirmed that the mismatch controls are equivalent to saline, as described in Example 19 below and in Table 1 of Nyce & Metzger, Nature 385, 721-725 (1997). Because of this finding, saline was used as a control for pulmonary function studies employing anti-sense oligos II, III and IV (SEQ. IS NOS; 7, 8 and 9).

Example 12: Specificity of Oligo I for Adenosine A₁ Receptor (Receptor Binding Studies)

Tissue from airway smooth muscle was dissected to primary, secondary and tertiary bronchi from rabbits which had been administered 20 mg oligo I (SEQ ID NO: 1) in 4 divided doses over a period of 48 hours as described above. A membrane fraction was prepared according to the method of Ali et al. (Ali, S., et al., Am. J. Resp. Crit. Care Med. 149: 908 (1994), the relevant section relating to the preparation of the membrane fraction is incorporated in its entirety hereby by reference).

The protein content was determined by the method of Bradford and plasma membranes were incubated with 0.2 U/ml adenosine deaminase for 30 minutes at 37°C to remove endogenous adenosine. See, Bradford, M. M. Anal. Biochem. 72, 240-254 (1976), the relevant portion of which is hereby incorporated in its entirety by reference. The binding of [3H]DPCPX, [3H]NPC17731, or [3H]CGS-21680 was measured as described by Jarvis et al. See, Jarvis, M.F., et al., Pharmacol. Exptl. Ther. 251, 888-893 (1989), the relevant portion of which is fully incorporated herein by reference. The results of this study are shown in Table 8 and discussed in Example 20 below.

25 Example 13: Pulmonary Function Measurements (Compliance c_{DYN} and Resistance)

At 4 months of age, the immunized animals were anesthetized and relaxed with 1.5 ml of a mixture of ketamine HCl (35 mg/kg) and acepromazine maleate (1.5 mg/kg) administered intramuscularly. After induction of anesthesia, allergic rabbits were comfortably positioned supine on a soft molded animal board. Salve was applied to the eyes to prevent drying, and they were closed. The animals were then intubated with a 4.0 mm intermediate high-low cuffed Murphy 1 endotracheal tube (Mallinckrodt, Glen Falls, NY), as previously described by Zavala and Rhodes. See, Zavala and Rhodes, Proc. Soc. Exp. Biol. Med. 144: 509-512

(1973), the relevant portion of which is incorporated herein by reference in its entirety. A polyethylene catheter of OD 2.4 mm (Becton Dickinson, Clay Adams, Parsippany NJ) with an attached thin-walled latex balloon was passed into the esophagus and maintained at the same distance (approximately 16 cm) from the mouth throughout the experiment. The endotracheal tube was attached to a heated Fleisch pneumotach (size 00; DEM Medical, Richmond, VA), and the flow (v) measured using a Validyne differential pressure transducer (Model DP-45-16-1927, Validyne Engineering, Northridge, CA), driven by a Gould carrier amplifier (Model 11-4113, Gould Electronics, Cleveland, OH).

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An esophageal balloon was attached to one side of the Validyne differential pressure transducer, and the other side was attached to the outflow of the endotracheal tube to obtain transpulmonary pressure (P_{tp}). The flow was integrated to yield a continuous tidal volume, and the measurements of total lung resistance (R_t) and dynamic compliance (C_{dyn}) were made at isovolumetric and zero flow points. The flow, volume and pressure were recorded on an eight channel Gould 2000 W high-frequency recorder and C_{dyn} was calculated using the total volume and the difference in P_{tp} at zero flow, and . R_t was calculated as the ratio of Ptp and V at midtidal lung volumes. These calculations were made automatically with the Buxco automated pulmonary mechanics respiratory analyzer (Model 6, Buxco Electronics, Sharon, CT), as previously described by Giles et al. See, Giles et al., Arch. Int. Pharmacodyn. Ther. 194: 213-232 (1971), the relevant portion of which describing these calculations is incorporated in toto hereby by reference. The results obtained upon administration of oligo II on allergic rabbits are shown and discussed in Example 26 below.

Example 14: Measurement of Bronchial Hyperresponsiveness (BHR)

Each allergic rabbit was administered histamine by aerosol to determine their baseline hyperresponsiveness. Aerosols of either saline or histamine were generated using a DeVilbiss nebulizer (DeVilbiss, Somerset, PA) for 30 seconds and then for 2 minutes at each dose employed. The ultrasonic nebulizer produced aerosol droplets of which 80% were < 5 micron in diameter. The histamine aerosol was administered in increasing concentrations (0.156 to 80 mg/ml) and measurements of pulmonary function were made after each dose. The B4R was then determined by calculating the concentration of histamine (mg/ml) required to reduce the $C_{\rm dyn}$ 50% from baseline (PC_{50 Histamine}).

Example 15: Cardiovascular Effect of Anti-sense Oligo I

The measurement of cardiac output and other cardiovascular parameters using Cardiomax™ utilizes the principal of thermal dilution in which the change in temperature of the blood exiting the heart after a venous injection of a known volume of cool saline is monitored. A single rapid injection of cool saline was made into the right atrium via cannulation of the

right jugular vein, and the corresponding changes in temperature of the mixed injectate and blood in the aortic arch were recorded via cannulation of the carotid artery by a temperature-sensing miniprobe.

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Twelve hours after the allergic rabbits had been treated with aerosols of oligo I (EPI 2010; SEQ. ID NO: 1) as described in (d) above, the animals were anesthetized with 0.3 ml/kg of 80% Ketamine and 20% Xylazine. This time point coincides with previous data showing efficacy for SEQ. ID NO: 1, as is clearly shown by Nyce & Metzger, (1997), supra, the pertinent disclosure being incorporated in its entirety here by reference. A thermocouple was then inserted into the left carotid artery of each rabbit, and was then advanced 6.5 cm and secured with a silk ligature. The right jugular vein was then cannulated and a length of polyethylene tubing was inserted and secured.

A thermodilution curve was then established on a CardiomaxTM II (Columbus Instruments, Ohio) by injecting sterile saline at 20°C to determine the correctness of positioning of the thermocouple probe. After establishing the correctness of the position of the thermocouple, the femoral artery and vein were isolated. The femoral vein was used as a portal for drug injections, and the femoral artery for blood pressure and heart rate measurements. Once constant baseline cardiovascular parameters were established, CardiomaxTM measurements of blood pressure, heart rate, cardiac output, total peripheral resistance, and cardiac contractility were made.

Example 16: Duration of Action of Oligo I (SEQ. ID NO: 1)

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Eight allergic rabbits received initially increasing log doses of adenosine by means of a nebulizer via an intra-tracheal tube as described in (f) above, beginning with 0.156 mg/ml until compliance was reduced by 50% ($PC_{50 \text{ Adenosine}}$) to establish a baseline. Six of the rabbits then received four 5 mg aerosolized doses of (SEQ. ID NO: 1) as described above. Two rabbits received equivalent amounts of saline vehicle as controls. Beginning 18 hours after the last treatment, the $PC_{50 \text{ Adenosine}}$ values were tested again. After this point, the measurements were continued for all animals each day, for up to 10 days. The results of this study are discussed in Example 25 below.

Example 17: Reduction of Adenosine A_{2b} Receptor Number by Anti-sense Oligo V

Sprague Dawley rats were administered 2.0 mg respirable anti-sense oligo V (SEQ ID NO:10) three times over two days using an inhalation chamber as described above. Twelve hours after the last administration, lung parenchymal tissue was dissected and assayed for adenosine A_{2b} receptor binding using [311]-NECA as described by Nyce & Metzger (1997), supra. Controls were conducted by administration of equal volumes of saline. The results are significant at p < 0.05 using Student's paired t test, and are discussed in Example 28 below.

Example 18: Comparison of Oligo I & Corresponding Phosphodiester Oligo VI (SEQ. ID NO:1681)

Oligo I (SEQ ID NO:1) countered the effects of adenosine and eliminated sensitivity to it for adenosine amounte up to 20 mg adenosine/5.0 ml (the limit of solubility of adenosine). Oligo VI (SEQ ID NO:1681), the phosphodiester version of the oligonucleotide sequence, was completely ineffective when tested in the same manner. Both compounds have identical sequence, differing only in the presence of phosphorothioate residues in Oligo I (SEQ ID NO:1), and were delivered as an aerosol as described above and in Nyce & Metzger (1997), supra. Significantly different at p<0.001, Student's paired t test. The results are discussed in Example 29 below..

RESULTS OBTAINED FOR ANTI-SENSE OLIGO I (SEQ. ID NO: 1)

Example 19: Results of Prior Work

The nucleotide sequence and other data for anti-sense oligo I (SEQ. ID NO: 1), which is

specific for the adenosine A₁ receptor, were provided above. The experimental data showing the effectiveness of oligo I in down regulating the receptor number and activity were also provided above.

Further information on the characteristics and activities of anti-sense oligo I is provided in Nyce, J. W. and Metzger, W. J., Nature 385:721 (1997), the relevant parts of which relating to the following results are incorporated in their entireties herein by reference. The Nyce & Metzger (1997) publication provided data showing that the anti-sense oligo I (SEQ. ID NO: 1):

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- (1) The anti-sense oligo I reduces the number of adenosine A₁ receptors in the bronchial smooth muscle of allergic rabbits in a dose-dependent manner as may be seen in Table 5 below.
- (2) Anti-sense Oligo I attenuates adenosine-induced bronchoconstriction and allergen-induced bronchoconstriction.
- (3) The Oligo I attenuates bronchial hyperresponsiveness as measured by PC_{50} histamine, a standard measurement to assess bronchial hyperresponsiveness. This result clearly demonstrates anti-inflammatory activity of the anti-sense oligo I as is shown in Table 5 above.
- (4) As expected, because it was designed to target it, the anti-sense oligo I is totally specific for the adenosine A_1 receptor, and has no effect at all at any dose on either the very closely related adenosine A_2 receptor or the related bradykinin B_2 receptor. This is seen in Table 5 below.
- (5) In contradistinction to the above effects of the Oligo I, the mismatch control molecules MM2 and MM3 (SEQ. ID NO:1682 and SEQ. ID NO:1683) which have identical base composition and molecular weight but differed from the anti-sense oligo I (SEQ ID NO: 1) by 6 and 2 mismatches, respectively. These mismatches, which are the minimum possible while still retaining identical base composition, produced absolutely no effect upon any of the targeted receptors (A₁, A₂ or B₂).

These results, along with a complete lack of prior art on the use of anti-sense oligonucleotides, such as oligo I, targeted to the adenosine A₁ receptor, are unexpected results. The showings presented in this patent clearly enable and demonstrate the effectiveness, for their intended use, of the claimed agents and method for treating a disease or condition associated with lung airway, such as bronchoconstriction, inflammation, allergy(ies), and the like.

Example 20: Oligo I Significantly Reduces Response to Adenosine Challenge

The receptor binding experiment is described in Example 12 above, and the results shown in Table 5 below which shows the binding characteristics of the adenosine A₁-selective ligand [3H]DPCPX and the bradykinin B₂-selective ligand [3H]NPC 17731 in membranes isolated from airway smooth muscle of A₁ adenosine receptor and B₂ bradykinin receptor anti-sense- and mismatch-treated allergic rabbits.

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Table 5: Binding Characteristics of Three Anti-Sense Oligos

	Treatment'	A ₁ r	eceptor	B ₂	receptor
	*	Kd	B _{max}	Kd	Bmax
	Adenosine A ₁	Receptor		· <u>-</u> · · · · · · · · · · · · · · · · · · ·	
	20 mg	0.36±0.029 nM	19±1.52 fmoles*	0.39±0.031 nM	14.8±0.99fmoles
5	2 mg	$0.38\pm0.030~\text{nM}$	32±2.56 fmoles*	0.41±0.028 nM	15.5 ± 1.08 fmoles
	0.2 mg	$0.37 \pm 0.030 \text{ nM}$	49±3.43 fmoles	0.34±0.024 nM	15.0±1.06 fmoles
	A ₁ MM1	(Control)			
	20 mg	0.34±0.027 nM	52.0±3.64 fmoles	0.35±0.024 nM	14.0±1.0 fmoles
	2 mg	0.37±0.033 nM	51.8±3.88 fmoles	0.38±0.028 nM	14.6±1.02 fmoles
10	B₂A (Bradykinin	Receptor)			
	20 mg	$0.36 \pm 0.028 \text{ nM}$	45.0±3.15 fmoles	0.38±0.027 nM	8.7±0.62 fmoles*
	2 mg	$0.39 \pm 0.035 \text{ nM}$	44.3 ± 2.90 fmoles	0.34±0.024 nM	11.9±0.76 fmoles**
	0.2 mg	0.40±0.028 nM	47.0±3.76 fmoles	0.35±0.028 nM	15.1±1.05 fmoles
	B ₂ MM (Control)				
15	20 mg	0.39±0.031 nM	42.0±2.94 fmoles	0.41 ±0.029 nM	14.0±0.98 fmoles
	2 mg	$0.41 \pm 0.035 \text{ nM}$	40.0±3.20 fmoles	$0.37 \pm 0.030 \text{ nM}$	14.8±0.99 fmoles
	0.2 mg	$0.37 \pm 0.029 \text{ nM}$	43.0±3.14 fmoles	$0.36 \pm 0.025 \text{ nM}$	15.1±1.35 fmoles
	Saline Control	0.37±0.041	46.0±5.21	0.39±0.047 nM	14.2±1.35 fmoles

Refers to total oligo administered in four equivalently divided doses over a 48 hour period. Treatments and analyses were performed as described in methods. Significance was determined by repeated-measures analysis of variance (ANOVA), and Tukey's protected t test. n = 4-6 for all groups.

* Significantly different from mismatch control- and saline-treated groups, p < 0.001;

**Significantly different from mismatch control- and saline-treated groups, p < 0.05.

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Example 21: Dose-response Effect of Oligo I

Anti-sense oligo I (SEQ ID NO:1) was found to reduce the effect of adenosine administration to the animal in a dose-dependent manner over the dose range tested as shown in Table 6 below.

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Table 6: Dose-Response Effect to Anti-sense Oligo I

Total Dose (mg)	PC ₅₀ Adenosine (mg Adenosine)
Anti-sense Oligo I	
0.2	8.32±7.2
2.0	14.0±7.2
20	19.5±0.34
A ₁ MM2 oligo (control)	
0.2	2.51 ± 0.46
2.0	3.13 ± 0.71
20	3.25 ± 0.34

The above results were studied with the Student's paired t test and found to be statistically different, p=0.05

The oligo I (SEQ. ID NO:1), an anti-adenosine A_1 receptor oligo, acts specifically on the adenosine A_1 receptor, but not on the adenosine A_2 receptors. These results stem from the treatment of rabbits with anti-sense oligo I (SEQ. ID NO.1) or mismatch control oligo (SEQ. ID NO:1682; A_1 MM2) as described in Example 9 above and in Nyce & Metzger (1997), supra (four doses of 5 mg spaced 8 to 12 hours apart via nebulizer via endotracheal tube), bronchial smooth muscle tissue excised and the number of adenosine A_1 and adenosine A_2 receptors determined as reported in Nyce & Metzger (1997), supra.

Example 22: Specificity of Oligo I (SEQ. ID NO:1) for Target Gene Product

Oligo I (SEQ. ID No:1) is specific for the adenosine A₁ receptor whereas its mismatch controls had no activity. Figure 1 depicts the results obtained from the cross-over experiment described in Example 10 above and in Nyce & Metzger (1997), supra. The two mismatch controls (SEQ. ID NO:1682 and SEQ. ID NO:1683) evidenced no effect on the PC_{50 Adenosine} value. On the contrary, the administration of anti-sense oligo I (SEQ. ID NO:1) showed a seven-fold increase in the PC_{50 Adenosine} value. The results clearly indicate that the anti-sense oligo I (SEQ. ID NO: 1) reduces the response (attenuates the sensitivity) to exogenously administered adenosine when compared with a saline control. The results provided in Table 2 above clearly establish that the effect of the anti-sense oligo I is dose dependent (see, column 3 of Table 1).

The Oligo I was also shown to be totally specific for the adenosine A_1 receptor, (see, top 3 rows of Table), inducing no activity at either the closely related adenosine A_2 receptor or the bradykinin B_2 receptor (see, lines 8-10 of Table 2 above).

In addition, the results shown in Table 2 establish that the anti-sense oligo I (SEQ. ID

NO:1) decreases sensitivity to adenosine in a dose dependent manner, and that it does this in an anti-sense oligo-dependent manner since neither of two mismatch control oligonucleotides (A_1MM2 ; SEQ. ID NO:1682 and A_1MM3 ; SEQ. ID NO:1683) show any effect on $PC_{50 \text{ Adenosine}}$ values or on attenuating the number of adenosine A_1 receptors.

5 Example 23: Effect on Aeroallergen-induced Bronchoconstriction & Inflammation

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The Oligo I (SEQ. ID NO:1) was shown to significantly reduce the histamine-induced effect in the rabbit model when compared to the mismatch oligos. The effect of the anti-sense Oligo I (SEQ. ID No:1) and the mismatch oligos (A₁MM2, SEQ. ID NO:1682 and A₁MM3, SEQ. ID NO:1682) on allergen-induced airway obstruction and bronchial hyperresponsiveness was assessed in allergic rabbits.

The effect of the anti-sense oligo I (SEQ. ID NO:1) on allergen-induced airway obstruction was assessed. As calculated from the area under the plotted curve, the anti-sense oligo I significantly inhibited allergen-induced airway obstruction when compared with the mismatched control (55%, p < 0.05; repeated measures ANOVA, and Tukey's t test).

A complete lack of effect was induced by the mismatch oligo A_iMM2 (Control) on allergen induced airway obstruction.

The effect of the anti-sense oligo I (SEQ. ID NO:1) on allergen-induced BHR was determined as above. As calculated from the $PC_{50 \; Histamine}$ value, the anti-sense oligo I (SEQ. ID NO:1) significantly inhibited allergen-induced BHR in allergic rabbits when compared to the mismatched control (61%, p<0.05; repeated measures ANOVA, Tukey's t test).

A complete lack of effect of the A₁MM mismatch control on allergen-induced BHR was observed.

The results indicated that anti-sense oligo I (SEQ. ID NO: 1) is effective to protect against aeroallergen-induced bronchoconstriction (house dust mite). In addition, the anti-sense oligo I (SEQ. ID NO:1) was also found to be a potent inhibitor of dust mite-induced bronchial hyper responsiveness, as shown by its effects upon histamine sensitivity which indicates anti-inflammatory activity for anti-sense oligo I (SEQ. ID NO:1).

Example 24: Anti-sense Oligo I is Free of Deleterious Side Effects

The Oligo I (SEQ. ID NO:1) was shown to be free of side effects that might be toxic to the recipient. No changes in arterial blood pressure, cardiac output, stroke volume, heart rate, total peripheral resistance or heart contractility (dPdT) were observed following administration of 2.0 or 20 mg oligo I (SEQ. ID NO:1). The addition, the results of the measurement of

cardiac output (CO), stroke volume (SV), mean arterial pressure (MAP), heart rate (HR), total peripheral resistance (TPR), and contractility (dPdT) with a Cardiomax[™] apparatus (Columbus Instruments, Ohio) were assessed.

These results evidenced that oligo I (SEQ. ID NO:1) has no detrimental effect upon critical cardiovascular parameters. More particularly, this oligo does not cause hypotension. This finding is of particular importance because other phosphorothioate anti-sense oligonucleotides have been shown in the past to induce hypotension in some model systems. Furthermore, the adenosine A₁ receptor plays an important role in sinoatrial conduction within the heart. Attenuation of the adenosine A₁ receptor by anti-sense oligo I (SEQ. ID NO:1) might be expected to result, therefore, in deleterious extrapulmonary activity in response to the downregulation of the receptor. This is not the case. The anti-sense oligo I (SEQ. ID NO:1) does not produce any deleterious intrapulmonary effects and renders the administration of the low doses of the present anti-sense oligo free of unexpected, undesirable side effects.

This demonstrates that when oligo I (SEQ. ID NO:1) is administered directly to the lung, it does not reach the heart in significant quantities to cause deleterious effects. This is in contrast to traditional adenosine receptor antagonists like theophylline which do escape the lung and can cause deleterious, even life-threatening effects outside the lung.

Example 25: Long Lasting Effect of Oligo I

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The Oligo I (SEQ. ID NO:1) evidenced a long lasting effect as evidenced by the PC_{50} and Resistance values obtained upon its administration prior to adenosine challenge.

The duration of the effect was measured for with respect to the PC_{50} of adenosine antisense oligo I when administered in four equal doses of 5 mg each by means of a nebulizer via an endotracheal tube, as described above. The effect of the agent is significant over days 1 to 8 after administration. When the effect of the anti-sense oligo I (SEQ. ID NO:1) had disappeared, the animals were administered saline aerosols (controls), and the $PC_{50 \text{ Adenosine}}$ values for all animals were measured again. Saline-treated animals showed base line $PC_{50 \text{ adenosine}}$ adenosine values (n=6).

The duration of the effect (with respect to Resistance) was measured for six allergic rabbits which were administered 20 mg of anti-sense oligo I (SEQ. ID NO: 1) as described above, upon airway resistance measured as also described above. The mean calculated duration of effect was 8.3 days for both PC_{50} adenosine (p<0.05) and resistance (p<0.05). These results show that anti-sense oligo I (SEQ. ID NO:1) has an extremely long duration of action, which is completely unexpected.

Example 26: Anti-sense Oligo II

Anti-sense oligo II, targeted to a different region of the adenosine A_i receptor mRNA, was found to be highly active against the adenosine A_i -mediated effects. The experiment measured the effect of the administration of anti-sense oligo II (SEQ. ID NO:7) upon compliance and resistance values when 20 mg anti-sense oligo II or saline (control) were administered to two groups of allergic rabbits as described above. Compliance and resistance values were measured following an administration of adenosine or saline as described above in Example 13. The effect of the anti-sense oligo of the invention was different from the control in a statistically significant manner, p < 0.05 using paired t-test, compliance; p < 0.01 for resistance.

The results showed that anti-sense oligo II (SEQ. ID NO:7), which targets the adenosine A_1 receptor, effectively maintains compliance and reduces resistance upon adenosine challenge.

Example 27: Antisense Oligos III and IV

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Oligos III (SEQ. ID NO:8) and IV (SEQ. ID NO:9) were shown to be in fact specifically targeted to the adenosine A₃ receptor by their effect on reducing inflammation and the number of inflammatory cells present upon separate administration of 20 mg of the anti-sense oligos III (SEQ. ID NO:8) and IV (SEQ. ID NO:9) to allergic rabbits as described above. The number of inflammatory cells was determined in their bronchial lavage fluid 3 hours later by counting at least 100 viable cells per lavage.

The effect of anti-sense oligos III (SEQ. ID NO:8) and IV (SEQ. ID NO:9) upon granulocytes, and upon total cells in bronchial lavage were assessed following exposure to dust mite allergen. The results showed that the anti-sense oligo IV (SEQ. ID NO:9) and antisense oligo III (SEQ. ID NO:8) are very potent anti-inflammatory agents in the asthmatic lung following exposure to dust mite allergen. As is known in the art, granulocytes, especially eosinophils, are the primary inflammatory cells of asthma, and the administration of anti-sense oligos III (SEQ. ID NO:8) and IV (SEQ. ID NO:9) reduced their numbers by 40% and 66%, respectively. Furthermore, anti-sense oligos IV (SEQ. ID NO:9) and III (SEQ. ID NO:8) also reduced the total number of cells in the bronchial lavage fluid by 40% and 80%, respectively. This is also an important indicator of anti-inflammatory activity by the present anti-adenosine A₃ agents of the invention. Inflammation is known to underlie bronchial hyperresponsiveness and allergen-induced bronchoconstriction in asthma. Both anti-sense oligonucleotides III (SEQ. ID NO:8) and IV (SEQ. ID NO:9), which are targeted to the adenosine A₃ receptor, are representative of an important new class of anti-inflammatory agents which may be designed to specifically target the lung receptors of each species.

Example 28: Anti-sense Oligo V

The anti-sense oligo V (SEQ. ID NO:10), targeted to the adenosine A_{2b} adenosine receptor mRNA was shown to be highly effective at countering adenosine A_{2b} -mediated effects

and at reducing the number of adenosine A_{2b} receptors present to less than half.

Example 29: Unexpected Superiority of Substituted over Phosphodiester-residue Oligo I-DS (SEQ. ID NO:1681)

Oligos I (SEQ. ID NO:1) and I-DS (SEQ. ID NO:1681) were separately administered to allergic rabbits as described above, and the rabbits were then challenged with adenosine. The phosphodiester oligo I-DS (SEQ. ID NO:1681) was statistically significantly less effective in countering the effect of adenosine whereas oligo I (SEQ. ID NO:1) showed high effectiveness, evidencing a PC_{50 Adenosine} of 20 mg.

10 Example 30: Anti-sense Oligo VI

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For the present work, I designed an additional anti-sense phosphorothioate oligo targeted to the adenosine A₁ receptor (Oligo VI). This anti-sense oligo was designed for therapy on a selected species as described in the above patent application and is generally specific for that species, unless the segment of the adenosine receptor mRNA of other species elected happens to have a similar sequence. The antisense oligos were prepared as described below, and tested in vivo in a rabbit model for bronchoconstriction, inflammation and lung allergy, which have breathing difficulties and impeded lung airways, as is the case in ailments such as asthma, as described in the above-identified application.

One additional oligo and its effect in a rabbit model was studied and the results of the study are reported and discussed below. The present oligo (anti-sense oligo VI) was selected for this study to complement the data on SEQ ID NO: 1 (Oligo I), which is anti-sense to the adenosine A₁ receptor mRNA provided in the above-identified patent application. This additional oligo is identified as anti-sense Oligo VI, and is targeted to a different region of the adenosine A₁ receptor mRNA than Oligo I. The design and synthesis of this anti-sense oligo was performed in accordance with the teaching, particularly Example 1, of the above-identified patent application.

The anti-sense Oligo VI is a phosphorothioate designed to target the coding region of the rabbit adenosine A_1 receptor mRNA region +964 to +984 relative to the initiation codon (start site). The Oligo VI was prepared as described in the above-indicated application, and is 20 nucleotides long. The OligoVI is directed to the adenosine A_1 receptor gene, and has the following sequence.

5'-CGC CGG CGG GTG CGG GCC GG-3' (SEQ. ID NO:_)

The phosphorothioate anti-sense Oligo VI having the sequence described in (5) above, was synthesized on an Applied Biosystems Model 396 Oligonucleotide Synthesizer, and purified using NENSORB chromatography (DuPont, DE). TETD (tetraethylthiuram disulfide) was used as the sulfurizing agent during the synthesis.

Example 31: Preparation of Allergic Rabbits

Neonatal New Zealand white Pasturella-free rabbits were immunized intraperitoneally within 24 hours of birth with 0.5 ml of 312 antigen units/ml house dust mite (D. farinae) extract (Berkeley Biologicals, Berkeley, CA) mixed with 10% kaolin as previously described (Metzger, W. J., in Late Phase Allergic Reactions, Dorsch, W., Ed., CRC Handbook, pp 347-362, CRC Press, Boca Raton,

1990; Ali, S. Et al., Am. J. Resp. Crit. Care Med. 149: 908 (1994)).

The immunizations were repeated weekly for the first month and then bi-weekly until the animals were 4 months old. These rabbits preferentially produce allergen-specific IgE antibody, typically respond to aeroallergen challenge with both an early and late-phase asthmatic response, and show bronchial hyper responsiveness (BHR). Monthly intraperitoneal administration of allergen (312 units dust mite allergen, as above) continues to stimulate and maintain allergen-specific IgE antibody and BHR. At 4 months of age, sensitized rabbits were prepared for aerosol administration as described by Ali et al. (1994), supra.

Example 32: Adenosine Aerosol Preparation

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An adenosine aerosol (20 mg/ml) was prepared with an ultrasonic nebulizer (Model 646, DeVilbiss, Somerset, PA), which produced aerosol droplets, 80% of which were smaller than 5μ m in diameter. Equal volumes of the aerosols were administered directly to the lungs via an intratracheal tube to all three rabbits.

The animals were then administered the aerosolized adenosine and Day 1 pre-treatment values for sensitivity to adenosine were calculated as the dose of adenosine causing a 50% loss of compliance (PC₅₀ Adenosine). The animals were then administered the aerosolized anti-sense via the intratracheal tube (5 mg/1.0 ml), for 2 minutes, twice daily for 2 days (total dose, 20 mg). Post-treatment PC₅₀ values were recorded (post-treatment challenge) on the morning of the third day. The results of these studies are provided in (9) below.

20 Example 33: Anti-sense Oligo Formulation

Each one of anti-sense oligos were separately solubilized in an aqueous solution and administered as described for anti-sense oligo I in (e) above, in four 5 mg aliquots (20 mg total dose) by means of a nebulizer via endotracheal tube, as described above.

Example 34: Oligo VI Reduces Response to Adenosine Challenge as well or Better than Oligo I

Oligo VI was tested in three allergic rabbits of the characteristics and readied as described in (7) above and in the above-indicated patent application. Oligo VI targets a section of the coding region of the A₁ receptor which is different from Oligo I. Both these target sequences were selected randomly from many possible coding region target sequences.

The three rabbits were treated identically as previously indicated for Oligo I. Briefly, 5 mg of Oligo VI were nebulized to the rabbits twice per day at 8 hour intervals, for two days. Thereafter, PC₅₀ adenosine studies were performed on the morning of the third day and compared to pre-treatment PC₅₀ values. This protocol is described in more detail in Nyce and Metzger (Nyce & Metzger, Nature 385: 721-725 (1997)).

The results obtained for the three rabbits are shown in Table 1 below.

Table 1:	PC ₅₀ Adenosine before & after Aerosolized Adenosine Treatment	
Treatment Time	PC ₅₀ Adenosine (mg)	

Pre-treatment 3.0 ± 2.1 Post-treatment>20.0*

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* maximum achievable dose due to adenosine insolubility in saline

All three animals treated with Oligo VI completely eliminated sensitivity to adenosine up to the measurable level of the agent shown in Table 1 above. That is, the administration of the Oligo VI abrogated the adenosine-induced bronchoconstriction in the three allergic rabbits. The actual efficacy of Oligo VI is, therefore, greater than could be measured in the experimental system used.

By comparing with the previously submitted results for the Oligo I, it may be seen that the Oligo VI was found to be as effective, or more, than Oligo I.

Example 34: Conclusions

The work described and results discussed in the examples clearly indicates that all antisense oligonucleotides designed in accordance with the teachings of the above-identified application were found to be highly effective at countering or reducing effects mediated by the receptors they are targeted to. That is, each and all of the two anti-sense oligos targeting an adenosine A₁ receptor mRNA, 1 anti-sense oligo targeting an adenosine A_{2b} receptor mRNA, and the 2 anti-sense oligos targeting an A₃ receptor mRNA were shown capable of countering the effect of exogenously administered adenosine which is mediated by the specific receptor they are targeted to.

The activity of the anti-sense oligos of this invention, moreover, is specific to the target and substitutively fails to inhibit another target.

In addition, the results presented also show that the administration of the present agents results in extremely low or non-existent deleterious side effects or toxicity.

This represents 100% success in providing agents that are highly effective and specific in the treatment of bronchoconstriction and/or inflammation. This invention is broadly applicable in the same manner to all gene(s) and corresponding mRNAs encoding proteins involved in or associated with airway diseases.

A comparison of the phosphodiester and a version of the same oligonucleotide wherein the phosphodiester bonds are substituted with phosphorothioate bonds evidenced an unexpected superiority for the phosphothiorate oligonucleotide over the phosphodiester anti-sense oligo.

The foregoing examples are illustrative of the present invention, and are not to be construed as limiting thereof. The invention is defined by the following claims, with equivalents of the claims to be included therein.

CLAIMS

1. An agent, comprising an oligonucleotide which is anti-sense to at least two mRNAs selected from the group consisting of target genes, coding and non-coding regions of RNAs corresponding to target genes, the genes' initiation codons, genomic flanking regions, intron-exon borders, the 5'-end, the 3'-end and the juxta-section between coding and non-coding regions and all segments of RNAs encoding proteins associated with one or more disease(s) or condition(s) or mixtures thereof.

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- The agent of claim 1, wherein one mRNA encodes a protein selected from the group consisting of transcription factors, stimulating and activating factors, interleukins, interleukin receptors, chemokines, chemokine receptors, endogenously produced specific and non-specific enzymes, immunoglobulins, antibody receptors, central nervous system (CNS) and peripheral nervous and non-nervous system receptors, CNS and peripheral nervous and non-nervous system peptide transmitters, adhesion molecules, defensines, growth factors, vasoactive peptides and receptors, and binding proteins, or corresponds to an oncogene.
- The agent of claim 2, wherein the encoded receptors and peptide transmitters selected from the group consisting of sympathomimetic receptors, parasympathetic receptors, GABA receptors, adenosine receptors, bradykinin receptors, insulin receptors, glucagon receptors, prostaglandin receptors, thyroid receptors, androgen receptors, anabolic receptors, estrogen receptors, progesterone receptors, receptors associated with the coagulation cascade, adenohypophyseal receptors, adenohypophyseal peptide transmitters, and histamine receptors (HisR).
- 4. The agent of claim 2, wherein the encoded sympathomimetic receptors and parasympathomimetic receptors are selected from the group consisting of acetylcholinesterase receptors (AcChaseR) acetylcholine receptors (AcChR), atropine receptors, muscarinic receptors, epinephrine receptors (EpiR), dopamine receptors (DOPAR), tachychinnen receptors, and norepinephrine receptors (NEpiR).
- 5. The agent of claim 2, wherein the encoded enzymes are selected from the group consisting of synthetases, kinases, oxidases, phosphatases, reductases, polysaccharide, triglyceride, and protein hydrolases, esterases, elastases, and, polysaccharide, triglyceride, lipid, and protein synthases.
- 6. The agent of claim 5, wherein the encoded enzymes are selected from the group consisting of tryptase, inducible nitric oxide synthase, cyclooxygenase (Cox), MAP kinase, eosinophil peroxidase, β2-adrenergic receptor kinase, leukotriene c-4 synthase, 5-lipoxygenase, phosphodiesterase IV, metalloproteinase, CSBP/p38 MAP kinase, neutrophil elastase, phospholipase A2, cyclooxygenase 2 (Cox-2), fucosyl transferase, IκB kinase 1 and 2, chymase, protein kinase C, thymidylate synthetase, tryptase, dihydrofolate reductase,

tryptase, thymidine kinase, deoxycytidine kinase, and ribonucleotide reductase.

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7. The agent of claim 2, wherein the encoded factor is selected from the group consisting of NfkB transcription factor, granulocyte macrophage colony stimulating factor (GM-CSF), AP-1 transcription factor, monocyte activating factor, neutrophil chemotactic factor, granulocyte colony-stimulating-factor (G-CSF), NFAT transcription factors, platelet activating factor, tumor necrosis factor α (TNF α), and basic fibroblast growth factor (BFGF).

- 8. The agent of claim 2, wherein the encoded adhesion molecule is selected from the group consisting of the intracellular adhesion molecules 1 (ICAM-1), 2 (ICAM-2) and 3 (ICAM-3), vascular cellular adhesion molecule (VCAM), endothelial leukocyte adhesion molecule-1 (ELAM-1), GATA transcription factor, neutrophil adherence receptor, mad CAM-1,
- 9. The agent of claim 2, wherein the encoded cytokines, lymphokines and chemokines are selected from the group consisting of the interleukin-1 (IL-1), interleukin-1 β (IL-1)interleukin-3 (IL-3), interleukin-4 (IL-4), interleukin-5 (IL-5), interleukin-8 (IL-8), interleukin-9 (IL-9), interleukin-11 (IL-11), CCR-5 CC chemokine, and Rantes.
- The agent of claim 2, wherein the encoded receptor is selected from the 10. group consisting of the adenosine A1 receptor, adenosine A2B receptor, adenosine A3 receptor, endothelin receptor A, endothellin receptor B, IgE high affinity receptor, muscarinic acetylcholine receptors, substance P receptor, histamine receptor, CCR-1 CC chemokine receptor, substance P, NK-1, and NK-3 receptors, CCR-2 CC chemokine receptor, CCR-3 CC chemokine receptor (Eotaxin Receptor) , interleukin-1 β receptor (IL-1 β R), interleukin-1 receptor (IL-1R), interleukin-1β receptor (IL-1βR), interleukin-3 receptor (IL-3R), CCR-4 CC chemokine receptor, cysteinyl leukotriene receptors, prostanoid receptors, interleukin-1 receptor (IL-1R), interleukin-4 receptor (IL-4R), interleukin-5 receptor (IL-5R), interleukin-8 receptor (IL-8R), interleukin-9 receptor (IL-9R), interleukin-11 receptor (IL-11R), bradykinin B2 receptor, sympathomimetic receptors, parasympathomimetic receptors, GABA receptors, adenosine receptors, bradykinin receptors, insulin receptors, glucagon receptors, prostaglandin receptors, thyroid receptors, androgen receptors, anabolic receptors, estrogen receptors, progesterone receptors, receptors associated with the coagulation cascade, adenohypophyseal receptors, and histamine receptors (HisR).
- 11. The agent of claim 2, wherein the encoded protein is selected from the group consisting of the eotaxin, major basic protein, preproendothelin, eosinophil cationic protein, P-selectin, STAT 4, STAT 6, c-mas, NF-Interleukin-6 (NF-IL-6), MIP-1α, MCP-2, MCP-3, MCP-4, cyclophillins, PDG2, cyclosporin A-binding protein, FK5-binding protein, fibronectin, LFA-1 (CD11a/CD18), PECAM-1, C3bi, PSGL-1,CD-34, substance P, p150,95, Mac-1 (CD11b/CD18), VLA-4, CD-18/CD11a, CD11b/CD18, C5a, CCR1, CCR2, CCR4, CCR5, and LTB-4.

12. The agent of claim 2, wherein the encoded defensin is selected from the group consisting of the defensin 1, defensin 2, and defensin 3.

13. The agent of claim 2, wherein the encoded selectin is selected from the group consisting of $\alpha 4\beta 1$ selectin, $\alpha 4\beta 7$ selectin, LFA-1 selectin, E-selectin, P-selectin, and L-selectin.

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- 14. The agent of claim 2, wherein the mRNA corresponds to an oncogene selected from the group consisting of ras, src, myc, and bcl-2.
- 15. The agent of claim 1, wherein at least one mononucleotide linking phosphodiester residue of the anti-sense oligonucleotide(s) is substituted by a residue selected from the group consisting of methylphosphonate, phosphotriester, phosphorothioate, phosphorodithioate, boranophosphate, formacetal, thioformacetal, thioether, carbonate, carbamate, sulfate, sulfonate, sulfamate, sulfonamide, sulfone, sulfite, sulfoxide, sulfide, hydroxylamine, methylene(methyimino), methyleneoxy (methylimino), 2'-O-methyl, phosphoramidate residues, and combinations thereof.
- 16. The agent of claim 15, wherein all phosphodiester residues are substituted.
- 17. The agent of claim 1, wherein the anti-sense oligonucleotide comprises about 7 to 60 mononucleotides.
- 18. The agent of claim 1, wherein the oligo consists of up to and including about 15% A.
 - 19. The agent of claim 1, wherein the oligo is adenosine-free.
- 20. The agent of claim 1, wherein the anti-sense oligo is selected from the group consisting of oligos comprising SEQ ID NOS.: __ through __, SEQ. ID NO:s__ to __ and SEQ. ID NOS: __ to __).
- The agent of claim 1, wherein at least one A is substituted by a universal base selected from the group consisting of heteroaromatic bases which bind to a thymidine base but have antagonist activity and less than about 0.3 of the adenosine base agonist activity at the adenosine A_1 , A_{2b} and A_3 receptors, and heteroaromatic bases which have no activity or have an agonist activity at the adenosine A_{2a} receptor.
- 22. The agent of claim 21, wherein the heteroaromatic bases are selected from the group consisting of pyrimidines and purines, which may be substituted by O, halo, NH₂, SH, SO, SO₂, SO₃, COOH and branched and fused primary and secondary amino, alkyl, alkenyl, alkynyl, cycloalkyl, heterocycloalkyl, aryl, heteroaryl, alkoxy, alkenoxy, acyl, cycloacyl, arylacyl, alkynoxy, cycloalkoxy, aroyl, arylthio, arylsulfoxyl, halocycloalkyl, alkylcycloalkyl, alkynylcycloalkyl, haloaryl, alkylaryl, alkenylaryl, alkynylaryl, arylalkyl, arylalkenyl, arylalkynyl, arylcycloalkyl, which may be further substituted by O, halo, NH₂, primary, secondary and tertiary amine, SH, SO, SO₂, SO₃, cycloalkyl, heterocycloalkyl and heteroaryl.

23. The agent of claim 22, wherein the pyrimidines and purines are substituted at positions 1, 2, 3, 4, 7 and 8.

24. The agent of claim 23, wherein the pyrimidines and purines are selected from the group consisting of theophylline, caffeine, dyphylline, etophylline, acephylline piperazine, bamifylline, enprofylline and xantine having the chemical formula

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- wherein R^1 and R^2 are independently H, alkyl, alkenyl or alkynyl and R^3 is H, aryl, dicycloalkyl, dicycloalkenyl, dicycloalkynyl, cycloalkyl, cycloalkenyl, cycloalkynyl, O-cycloalkynyl, NH₂-alkylamino-ketoxyalkyloxy-aryl and mono and dialkylaminoalkyl-N-alkylamino-SO₂ aryl.
- 25. The agent of claim 24, wherein the universal base is selected from the group consisting of 3-nitropyrrole-2'-deoxynucleoside, 5-nitro-indole, 2-deoxyribosyl-(5-nitroindole), 2-deoxyribofuranosyl-(5-nitroindole), 2'-deoxynosine, 2'-deoxynebularine, 6H, 8H-3,4-dihydropyrimido [4,5-c] oxazine-7-one or 2-amino-6-methoxyaminopurine.
- 26. The agent of claim 1, where a methylated cytocine (ⁿC) is substituted for at least one CpG dinucleotide if present in the oligo(s).
- 27. The agent of claim 1, wherein the anti-sense oligonucleotide is operatively linked to a cell internalized or up-taken agent or to a eukaryotic or prokaryotic vector.
- 28. The agent of claim 27, wherein the cell internalized or up taken agent is selected from the group consisting of transferrin, asialoglycoprotein, and streptavidin.
- 29. A composition, comprising the agent of claim 1 and a pharmaceutically or veterinarily acceptable carrier.
- 30. The composition of claim 29, wherein the carrier is selected from the group consisting of gaseous, liquid and solid carriers.
- 31. The composition of claim 29, further comprising an agent selected from the group consisting of other therapeutic compounds, surfactants, antioxidants, flavoring and coloring agents, fillers, volatile oils, buffering agents, dispersants, RNA inactivating agents, antioxidants, flavoring agents, propellants and preservatives.
- 32. The composition of claim 29, wherein the anti-sense oligonucleotide is present in an amount of about 0.01 to about 99.99 w/w of the composition.
 - 33. The composition of claim 32, comprising the nucleic acid, a surfactant and a carrier.
 - 34. The composition of claim 33, wherein the surfactant is selected from the

group consisting of surfactant protein A, surfactant protein B, surfactant protein C, surfactant protein D and surfactant protein and active fragments thereof, non-dipalmitoyl disaturated phosphatidylcholine, dipalmitoylphosphatidylcholine, phosphatidylcholine, phosphatidylglycerol, phosphatidylinositol, phosphatidylethanolamine, phosphatidylserine, phosphatidic acid, ubiquinones, lysophosphatidylethanolamine, lysophosphatidylcholine, palmitoyl-lysophosphatidylcholin, dehydroepiandrosterone, dolichols, sulfatidic acid, glycerol-3-phosphate, dihydroxyacetone phosphate, glycerol, glycero-3-phosphocholine, dihydroxyacetone, palmitate, cytidine diphosphate (CDP) diacylglycerol, CDP choline, choline, choline phosphate, artificial lamellar bodies vehicles for surfactant components, omega-3 fatty acids, polyenic acid, polyenoic acid, lecithin, palmitic acid, non-ionic ethylene and/or propylene oxide block copolymers, polyoxypropylene, polyoxyethylene, poly (vinyl amine) with dextran and/or alkanoyl side chains, Brij 35, Triton X-100, ALEC, Exosurf, Survant and Atovaquone.

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- 35. A formulation, comprising the composition of claim 26, wherein the carrier comprises a hydrophobic carrier.
- 36. The formulation of claim 35, wherein the carrier comprises lipid particles or vesicles.
- 37. The formulation of claim 36, wherein the vesicles comprise liposomes and the particles comprise micro crystals.
- 38. The formulation of claim 35, wherein the lipid vesicles comprise N-(1-[2, 3-dioleoxyloxi] propyl) -N,N,N- trimethyl- ammonium methylsulfate.
 - 39. The formulation of claim 34, comprising a respirable formulation.
 - 40. The formulation of claim 34, comprising an aerosol.
 - 41. The formulation of claim 34, in single or multiple unit form.
 - 42. The formulation of claim 34, in bulk.
 - 43. A capsule or cartridge, comprising the composition of claim 26.
- 44. A kit, comprising a delivery device, in a separate container, the composition of claim 26 and instructions for its use.
- 45. The kit of claim 44, wherein the delivery device comprises a nebulizer which delivers single metered doses of the formulation.
- 46. The kit of claim 44, wherein the nebulizer comprises an insufflator, and the composition is provided in a piercable or openable capsule or cartridge.
- 47. The kit of claim 44, wherein the delivery device comprises a pressurized inhaler, and the composition comprises a suspension or solution of the agent.
- 48. The kit of claim 44, further comprising in a separate container an agent selected from the group consisting of other therapeutic compounds, surfactants, antioxidants, flavoring and coloring agents, fillers, volatile oils, buffering agents, dispersants, cell internalized or up taken agents, RNA inactivating agents, antioxidants, flavoring agents,

propellants and preservatives.

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- The kit of claim 48, wherein the surfactant is selected from the group 49. consisting of surfactant protein A, surfactant protein B, surfactant protein C, surfactant protein D and surfactant protein and active fragments thereof, non-dipalmitoyl disaturated phosphatidylcholine, dipalmitoylphosphatidylcholine, phosphatidylcholine, phosphatidylglycerol, phosphatidylinositol, phosphatidylethanolamine, phosphatidylserine, phosphatidic acid, ubiquinones, lysophosphatidylethanolamine, lysophosphatidylcholine, palmitoyl-lysophosphatidylcholin, dehydroepiandrosterone, dolichols, sulfatidic acid, glycerol-3-phosphate, dihydroxyacetone phosphate, glycerol, glycero-3-phosphocholine, dihydroxyacetone, palmitate, cytidine diphosphate (CDP) diacylglycerol, CDP choline, choline, choline phosphate, artificial lamellar bodies vehicles for surfactant components, omega-3 fatty acids, polyenic acid, polyenoic acid, lecithin, palmitic acid, non-ionic ethylene and/or propylene oxide block copolymers, polyoxypropylene, polyoxyethylene, poly (vinyl amine) with dextran and/or alkanoyl side chains, Brij 35, Triton X-100, ALEC, Exosurf, Survant and Atovaquone.
- 50. The kit of claim 44, wherein the composition is provided in a capsule or cartridge.
 - 51. A cell, comprising the agent of claim 1.
- 52. A method of treating a disease or condition associated with the mRNA corresponding to at least one target gene(s), genomic flanking regions, or proteins, comprising administering to a subject afflicted with the disease or condition the agent of claim 1, comprising an amount of the anti-sense oligonucleotide effective to reduce the production or availability, or to increase the degradation by the subject of at least one of the target mRNA.
- 53. The method claim 52, wherein the agent is administered in an amount effective to reduce the production or availability, or to increase the degradation of at least two of the target mRNAs.
 - 54. The method of claim 52, wherein the agent is administered directly to the lung (s) of the subject.
- 55. The method of claim 54, wherein the agent is administered as a respirable aerosol.
 - 56. The method of claim 52, wherein the disease or condition is a lung disease or condition, and at least one of the target mRNA encodes a protein selected from the group consisting of the adenosine A_1 receptor, adenosine A_2 B receptor, adenosine A_3 receptor, and bradykinin B2 receptor.
 - 57. The method of claim 56, wherein the disease or condition is associated with obstruction of the subject's airways.
 - 58. The method of claim 57, wherein the disease or condition is associated with asthma.

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/19419

A. CLASSIFICATION OF SUBJECT MATTER IPC(6): A61K 31/70, 48/00; C07H 21/00, 21/04; C12N 5/10 US CL: 435/325, 375; 514/44; 536/24.5 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) U.S.: 435/325, 375; 514/44; 536/24.5 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) Please Sec Extra Sheet.			
C. DOCUMENTS CONSIDERED TO BE RE			
Category* Citation of document, with indicat	on, where appropriate, of the relevant passages Relevant to claim No.		
abstract, page 4, line 19, to	3, 4, 6-16, 18-26, 28-50, 54-58		
Y,P WO 98/23294 A1 (EAST of 1998, see page 3, line 23, to bridging sentence.	page 29, line 24, especially pages 3-4,		
Further documents are listed in the continua	tion of Box C. See patent family annex.		
Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "B" cartier document published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is			
cited to establish the publication date of another of special reason (as specified) "O" document referring to an oral disclosure, use, exhimeans	considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art		
P document published prior to the international filing da the priority date claimed			
Date of the actual completion of the international of DECEMBER 1998	Date of mailing of the international search report 11 JAN 1999		
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. (703) 305-3230 Authorized officer Journal Total THOMAS G. LARSON, Ph.D. Telephone No. (703) 308-0196			

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/19419

found unsearchable (Continuation of item 1 of first sheet)		
Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)		
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:		
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:		
·		
2. X Claims Nos.: 20, 24, and 25 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:		
Please See Extra Sheet.		
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).		
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)		
This International Searching Authority found multiple inventions in this international application, as follows:		
,		
·		
N annual N annual All		
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.		
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.		
The specific of the specific o		
As only some of the required additional search lees were timely paid by the application and only those claims for which fees were paid, specifically claims Nos.:		
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:		
Remark on Protest The additional search fees were accompanied by the applicant's protest.		
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.		

Form PCT/ISA/210 (continuation of first sheet(1))(July 1992)*

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/19419

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, STN (BIOSIS, CAPLUS, INPADOC, LIFESCI, MEDLINE, WPIDS)
Search Terms: antisense, multitarget, multi, multiple, two, targets, asthma, transcripts, receptor, synthetase, synthase, phosphatase, esterase, Nyce, Jonathan.

BOX I. OBSERVATIONS WHERE CLAIMS WERE FOUND UNSEARCHABLE

2. Where no meaningful search could be carried out, specifically:

Claim 20 could not be search because the SEQ. ID. NOS. are missing from the claim and because a copy of the sequence listing in computer readable format (CRF) was not provided (see PCT/RO/101, Box No. VIII, item no. 8). Claim 24 and dependent claim 25 could not be examined because claim 24 is missing the chemical formula recited in the claim.